PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:		(11) International Publication Number: WO 95/15381
C12N 15/12, C12Q 1/68, A01K 67/00, A61K 48/00, C07K 14/47	A2	(43) International Publication Date: 8 June 1995 (08,06,95)
(21) International Application Number: PCT/US	94/138	05 (81) Designated States: CA, JP, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).
(22) International Filing Date: 2 December 1994 (02.12.9	04)
(30) Priority Data: 160,295 2 December 1993 (02.12.93) (Published Without international search report and to be republished upon receipt of that report.
(71) Applicant: THE JOHNS HOPKINS UNIVERSITY 720 Rutland Avenue, Baltimore, MD 21205 (US).		sj;
(71)(72) Applicant and Inventor: DE LA CHAPELLE [FVFI]; Bregmansgatan 11, FIN-00140 Helsingford		еп
(72) Inventors: VOGELSTEIN, Bert; 3700 Breton Way, B MD 21208 (US). KINZLER, Kenneth, W.; 1348 Road, Baltimore, MD 21234 (US).		
(74) Agents: KAGAN, Sarah, A. et al.; Banner, Birch, I Beckett, 11th floor, 1001 G Street, N.W., Washin 20001 (US).		
(54) Title: HUMAN MUTATOR GENE hMSH2 AND H	ERED	ITARY NON POLYPOSIS COLORECTAL CANCER
(57) Abstract		
MuiS class of genes, which are involved in DNA mismate	ch repa exister	lyposis colorectal cancer, was identified by virtue of its homology to the lir. The sequence of cDNA clones of the human gene are provided, and nee of germ line mutations in hereditary non-polyposis colorectal cancer r cells.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

					•
AT	Austria	GB	United Kingdom	MR	Mauritania
AU	Australia	GE	Georgia	MW	Malawi
BB	Barbados	GN	Guinea	NE	Niger
BE	Belgium	GR	Greece	NL	Netherlands
BF	Burkina Faso	HU	Hungary	NO	Norway
BG	Bulgaria	1E	Ireland	NZ	New Zealand
BJ	Benin	IT	Italy	PL	Poland
BR	Brazil	JP	Japan	PT	Portugal
BY	Belarus	KE	Kenya	RO	Romania
CA	Canada	KG	Kyrgystan	RU	Russian Federation
CF	Central African Republic	KP	Democratic People's Republic	\$D	Sudan
CG	Congo		of Korea	SE	Sweden
CH	Switzerland	KR	Republic of Korea	SI	Slovenia
CI	Côte d'Ivoire	ΚZ	Kazakhstan	SK	Slovakia
CM	Cameroon	LI	Liechtenstein	SN	Senegal
CN	China	LK	Sri Lanka	TD	Chad
CS	Czechoslovakia	LU	Luxembourg	TG	Togo
CZ	Czech Republic	LV	Latvia	LT	Tajikistan
DE	Germany	MC	Monaco	TT	Trinidad and Tobago
DK	Denmark	MĐ	Republic of Moldova	UA	Ukraine
E.S	Spain	MG	Madagascar	US	United States of America
Fī	Finland	ML	Mali	UZ	Uzbekistan
FR	France	MN	Mongolia	VN	Viet Nam
C.A	Cahon		•		

HUMAN MUTATOR GENE hMSH2 AND HEREDITARY NON POLYPOSIS COLORECTAL CANCER

This invention was made using U.S. government grants from the NIH CA47527, CA09320, GM26449, CA09243, CA41183, CA42705 CA57435, and CA35494, as well as grants from the Department of Energy DOE/ERN/F139 and DE-FG 09291ER-61139. Therefore the U.S. government retains certain rights to the invention.

This application is a continuation-in-part of Serial No. 08/056,546, filed May 5, 1993.

TECHNICAL FIELD OF THE INVENTION

The invention relates to a gene which predisposes individuals to colorectal and other cancers. In addition, it also relates to biochemical tests which can be used to identify drugs for treatment of affected individuals.

BACKGROUND OF THE INVENTION

HNPCC (Lynch syndrome) is one of the most common cancer predisposition syndromes, affecting as many as 1 in 200 individuals in the western world (Lynch et al., 1993). Affected individuals develop tumors of the colon, endometrium, ovary and other organs, often before 50 years of age. Although the familial nature of this syndrome was discovered nearly a century ago (Warthin et al., 1913), the role of heredity in its causation remained difficult to define (Lynch et al., 1966). Recently, however, linkage analysis in two large kindreds demonstrated association with polymorphic markers on chromosome 2 (Peltomaki

WO 95/15381

et al., 1993a). Studies in other families suggested that neoplasia in a major fraction of HNPCC kindreds is linked to this same chromosome 2p locus (Aaltonen et al., 1993).

HNPCC is defined clinically by the occurrence of early-onset colon and other specific cancers in first degree relatives spanning at least two generations (Lynch et al., 1993). The predisposition is inherited in an autosomal dominant fashion. It was initially expected that the gene(s) responsible for HNPCC was a tumor suppressor gene, as other previously characterized cancer predisposition syndromes with this mode of inheritance are caused by suppressor gene mutations (reviewed in Knudson, 1993). But the analysis of tumors from HNPCC patients suggested a different mechanism. Most loci encoding tumor suppressor genes undergo somatic losses during tumorigenesis (Stanbridge, 1990). In contrast, both alleles of chromosome 2p loci were found to be retained in HNPCC tumors (Aaltonen et al., 1993). During this search for chromosome 2 losses, however, it was noted that HNPCC tumors exhibited somatic alterations of numerous microsatellite sequences.

Widespread, subtle alterations of the cancer cell genome were first detected in a subset of sporadic colorectal tumors using the arbitrarily-primed polymerase chain reaction (Peinado et al., 1992). These alterations were subsequently found to represent deletions of up to 4 nucleotides in genomic polyA tracts (Ionov et al., 1993). Other studies showed that a similar, distinctive subgroup of sporadic tumors had insertions or deletions in a variety of simple repeated sequences, particularly microsatellite sequences consisting of dinucleotide or trinucleotide repeats (Ionov et al., 1993; Thibodeau et al., 1993; Aaltonen et al., 1993). Interestingly, these sporadic tumors had certain features in common with those developing in HNPCC kindreds, such as a tendency to be located on the right side of the colon and to be near-diploid. These and other data suggested that HNPCC and a subset of sporadic tumors were associated with a heritable defect causing replication errors (RER) of microsatellites (Ionov et al., 1993; Aaltonen et al., 1993).

The mechanism underlying the postulated defect could not be determined from the study of tumor DNA, but studies in simpler organisms provided an intriguing possibility (Levinson and Gutman, 1987; Strand et al., 1993). This work showed that bacteria and yeast containing defective mismatch repair genes manifest instability of dinucleotide repeats. The disruption of genes primarily involved in DNA replication or recombination had no apparent effect on the fidelity of microsatellite replication (reviewed in Kunkel, 1993). These pivotal studies suggested that defective mismatch repair might be responsible for the microsatellite alterations in the tumors from HNPCC patients (Strand et al., 1993).

Thus there is a need in the art to identify the actual gene and protein responsible for hereditary non-polyposis colorectal cancer and the replication error phenotype found in both hereditary and sporadic tumors. Identification of the gene and protein would allow more widespread diagnostic screening for hereditary non-polyposis colorectal cancer than is currently possible. Identification of the involved gene and protein would also enable the rational screening of compounds for use in drug therapy of hereditary non-polyposis colorectal cancer, and would enable gene therapy for affected individuals.

SUMMARY OF THE INVENTION

It is an object of the invention to provide a DNA molecule which when mutated is the genetic determinant for hereditary non-polyposis colorectal cancer.

It is another object of the invention to provide DNA molecules which contain specific mutations which cause hereditary non-polyposis colorectal cancer.

It is yet another object of the invention to provide methods of treating persons who are predisposed to hereditary non-polyposis colorectal cancer.

It is still another object of the invention to provide methods for determining a predisposition to cancer.

It is a further object of the invention to provide methods for screening test compounds to identify therapeutic agents for treating persons predisposed to hereditary non-polyposis colorectal cancer.

It is still another object of the invention to provide a protein which is important for human DNA mismatch repair.

It is yet another object of the invention to provide a transgenic animal for studying potential therapies for hereditary non-polyposis colorectal cancer.

These and other objects of the invention are provided by one or more of the embodiments described below. In one embodiment of the invention an isolated and purified DNA molecule is provided. The molecule has a sequence of at least about 20 nucleotides of hMSH2, as shown in SEQ ID NO:1.

In another embodiment of the invention an isolated and purified DNA molecule is provided. The DNA molecule has a sequence of at least about 20 nucleotides of an hMSH2 allele found in a tumor wherein said DNA molecule contains a mutation relative to hMSH2 shown in SEQ ID NO:1.

In yet another embodiment of the invention a method of treating a person predisposed to hereditary non-polyposis colorectal cancer is provided. The method prevents accumulation of somatic mutations. The method involves administering a DNA molecule which has a sequence of at least about 20 nucleotides of hMSH2, as shown in SEQ ID NO:1, to a person having a mutation in an hMSH2 allele which predisposes the person to hereditary non-polyposis colorectal cancer, wherein said DNA molecule is sufficient to remedy the mutation in an hMSH2 allele of the person.

In another embodiment of the invention a method is provided for determining a predisposition to cancer. The method involves testing a body sample of a human to ascertain the presence of a mutation in hMSH2 which affects hMSH2 expression or hMSH2 protein function, the presence of such a mutation indicating a predisposition to cancer.

In still another embodiment of the invention a method is provided for screening to identify therapeutic agents which can prevent or ameliorate tumors. The screening method involves contacting a test compound with a purified hMSH2 protein or a cell; determining the ability of the hMSH2 protein or the cell to perform DNA mismatch repair, a test compound which increases the ability of said

hMSH2 protein or said cell to perform DNA mismatch repair being a potential therapeutic agent.

In another embodiment of the invention an isolated and purified protein is provided. The protein has the sequence shown in SEQ ID NO:2.

In still another embodiment of the invention a transgenic animal is provided. The transgenic (nonhuman) animal maintains an hMSH2 allele in its germline. The hMSH2 allele is one which is found in humans having hereditary non-polyposis colorectal cancer or in RER⁺ tumors. Also provided are animals which have no wild-type MSH2 alleles, due to mutations introduced.

Thus the present invention provides the art with the sequence of the gene responsible for hereditary non-polyposis colorectal cancer and information regarding the mechanism by which it causes tumors. This enables the art to practice a variety of techniques to identify persons at risk of developing a variety of cancers and to treat them to prevent such cancers from actually developing.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 summarizes the markers retained in somatic cell hybrids used in locating the hMSH2 gene.

PCR was used to determine whether each of the listed markers was present (black box) or absent (white box)in the indicated hybrid. The laboratory name of each hybrid and the formal name (in parentheses) is listed. The hybrid panel was also validated with ten additional polymorphic markers outside of the 136-177 region. M: hybrid derived from microcell-mediated chromosome 2 transfer; T: derived from t(X;2) translocation; X: derived from X-irradiated chromosome 2 donor. MO: mouse-human hybrid; HA: hamster-human hybrid; RA: rat-human hybrid; TEL: telomere; CEN: centromere.

Figure 2 shows a FISH analysis which was used to determine the proximity and ordering of DNA sequences within chromosome band 2p16.

Panels 2A and 2B show FISH mapping of the 123 marker. Panel 2A shows G-banded metaphase chromosome 2. Panel 2B shows identical chromosome as in

Panel 2A following FISH with a biotin-labeled P1 clone for the 123 marker. Results localize the 123 marker to chromosome band 2p16.3. Panels 2C and 2D show co-hybridization documenting the coincident localization of a microdissection (Micro-FISH) probe from chromosome 2p16 and the 123 marker. Panel 2C shows DAPI stained metaphase chromosome 2. Panel 2D shows simultaneous hybridization of the biotin-labeled 123 probe (appearing as an intensely staining smaller circle) and the Spectrum-Orange labeled 2p16 Micro-FISH probe (appearing as a diffusely staining larger circle). Panel 2E shows a representative example of an interphase nucleus simultaneously hybridized with P1 clones for CA5, hMSH2 and ze3. The results were used to directly measure the distances between markers in order to establish the order and relative distance between markers (after Trask et al., 1989). Inset: The image processing program NIH Image was used to provide an average gray value displayed as a surface plot to support the length measurements and to graphically illustrate the relative order The surface plot presented defines the specified interphase chromosome and the relative order CA-MSH2-ze3.

Figure 3 shows linkage analysis of HNPCC pedigrees.

All affected individuals in which meiotic recombination occurred between markers 119 and 136 are included. A black box indicates that the individual did not contain the allele associated with disease in his/her family or that the individual inherited an allele not associated with disease from his/her affected parent. A white box indicates that the individual had an allele which was the same size as the disease-associated allele. A hatched circle indicates that the marker was not studied. All individuals had colon or endometrial cancer at less than 55 years of age, or had progeny with such disease but did not indicate that the patient necessarily had disease-associated alleles because phase could usually not be determined.

Figure 4 shows hMSH2 gene localization.

Southern blots containing EcoRI (figures 4A and 4C) and PstI (figures 4B and 4D) digested DNA from the indicated somatic cell hybrids (figures 4A and 4B)

or YAC clones (figures 4C and 4D) were hybridized with a radiolabelled insert from cDNA clone pNP-23. Southern blotting and hybridization were performed as described (Vogelstein et al., 1987). Autoradiographs are shown. The 5.0 kb PstI fragment in hybrids Z11 and Z12 is derived from hamster DNA.

Figure 5 shows the cDNA sequence of hMSH2.

An open reading frame (ORF) begins at nucleotide 1 and ends at nt 2802. The predicted amino acid sequence is shown. The sequence downstream of nt 2879 was not determined.

Figure 6 shows homology between yeast and human MSH2 genes.

The predicted amino acid sequences of yeast (y) MSH2 (Reenan and Kolodner, 1992) and human MSH2 genes are compared within the region of highest homology. Blocks of similar amino acids are shaded.

Figure 7 shows germline and somatic mutations of hMSH2.

Autoradiographs of polyacrylamide gels containing the sequencing reactions derived from PCR products are shown. The 1.4 kb PCR products containing a conserved region of hMSH2 were generated from genomic DNA samples as described in the Examples. Antisense primers were used in the sequencing reactions. The ddA mixes from each sequencing reaction were loaded in adjacent lanes to facilitate comparison, as were those for C, G, and T. The DNA samples were derived from the tumor (lane 1) and normal colon (lane 2) of patient Cx10, an RER- colon tumor cell line (lane 3), and lymphocytes of patients J-42 (lane 4) and J-143 (lane 5). Figure 7A: A transition (C to T at codon 622) in lymphocyte DNA can be observed in HNPCC patients J-42 and J-143. Figure 7B: A transition (C to T at nt codon 639) in tumor (lane 1) and normal colonic mucosa (lane 2) of patient Cx10. Figure 7C: A substitution of a TG dinucleotide for an A at codon 663 can be observed in DNA of the tumor of patient Cx10, (lane 1), but not in DNA from her normal colon (lane 2). Arrows mark the substitutions in panel A and B and the TG dinucleotide insertion site in panel C.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

The disclosure of Serial No. 08/056,546, filed May 5, 1993, is expressly incorporated herein.

It is a discovery of the present invention that the gene responsible for hereditary non-polyposis colorectal cancer is hMSH2, a human analog of bacterial MutS. The cDNA sequence of hMSH2 is shown in SEQ ID NO:1. This gene encodes a DNA mismatch repair enzyme. Mutation of the gene causes cells to accumulate mutations. For example, the observed replication error phenotype (RER+) found in both sporadic and hereditary non-polyposis colorectal cancer consists of variations (insertions and deletions) in microsatellite DNA. In yeast and bacteria defective MutS-related genes cause other types of mutations as well.

Useful DNA molecules according to the invention are those which will specifically hybridize to hMSH2 sequences. Typically these are at least about 20 nucleotides in length and have the nucleotide sequence as shown in SEQ ID NO:1. Such molecules can be labeled, according to any technique known in the art, such as with radiolabels, fluorescent labels, enzymatic labels, sequence tags, etc. According to another aspect of the invention, the DNA molecules contain a mutation which has been found in tumors of HNPCC patients or in sporadic RER⁺ tumors. Such molecules can be used as allele-specific oligonucleotide probes to track a particular mutation through a family.

According to some aspects of the invention, it is desirable that the DNA encode all or a part of the hMSH2 protein as shown in SEQ ID NO:2. To obtain expression of the protein the DNA sequence can be operably linked to appropriate control sequences, such as promotor, Kozak consensus, and terminator sequences.

A person who is predisposed to develop cancers due to inheritance of a mutant hMSH2 allele can be treated by administration of a DNA molecule which contains all or a part of the normal hMSH2 gene sequence as shown in SEQ ID NO:1. A portion of the gene sequence will be useful when it spans the location of the mutation which is present in the mutant allele, so that a double recombination event between the mutant allele and the normal portion "corrects"

the defect present in the person. A portion of the gene can also be usefully administered when it encodes enough of the protein to express a functional DNA mismatch repair enzyme. Such a portion need not necessarily recombine with the mutant allele, but can be maintained at a separate locus in the genome or on an independently replicating vector. Means for administering DNA to humans are known in the art, and any can be used as is convenient. A variety of vectors are also known for this purpose. According to some techniques vectors are not required. Such techniques are well known to those of skill in the art.

Also contemplated as part of the present invention is the use of a combined anti-neoplastic therapy regimen. Such a combined regimen is useful for patients having an RER+ tumor, whether sporadic or associated with HNPCC. The regimen combines any standard anti-neoplastic therapy to which a patient can become resistant and hMSH2 gene therapy, as described above. By remedying the defect present in RER+ cells, i.e., an hMSH2 mutation, the likelihood of the tumor developing a resistance mutation is greatly diminished. By delaying or preventing the onset of resistance, the life of cancer patients can be prolonged. In addition, such prevention of resistance allows a greater degree of tumor destruction by the therapeutic agent. Examples of anti-neoplastic therapies which can be combined with hMSH2 gene therapy are hormones, radiation, cytotoxic drugs, cytotoxins, and antibodies.

Body samples can be tested to determine whether the hMSH2 gene is normal or mutant. Mutations are those deviations from the sequence shown in SEQ ID NO:1 which are associated with disease and which cause a change in hMSH2 protein function or expression. Such mutations include nonconservative amino acid substitutions, deletions, premature terminations and frameshifts. See Table I. Suitable body samples for testing include those comprising DNA, RNA, or protein, obtained from biopsies, blood, prenatal, or embryonic tissues, for example.

Provided with the information that the defect causing HNPCC and sporadic RER⁺ tumors is in a DNA mismatch repair enzyme, one can perform assays on

test compounds and compositions to determine if they will remedy the defect. Such therapeutic compounds could bind to missense hMSH mutant proteins to restore the proteins to the normal, active conformation. Alternatively such therapeutic compounds could stimulate the expression of alternate pathways for mismatch repair. Screening for such therapeutic compounds could be performed by contacting test compounds with cells, either normal or those with an hMSH2 mutation found in a tumor. The ability of the cells which were contacted with the test compounds is compared with the ability of the same cells which were not contacted for mismatch repair activity. Such activity can be tested as is known in the art. See, for example, Levinson and Gutman, 1987, and Strand et al., 1993. Observation of changes in microsatellite DNA in cells is one way of assessing mismatch repair activity. Another approach is to assay DNA mismatch repair in vitro in nuclear extracts. See Holmes, 1990; Thomas, 1991; and Fang, 1993.

Fable I

Sample	Source	Type	Codon	cDNA Nucleotide Change	Predicted Coding Change
Family J	HNPCC Kindred	Germline	622	CCA to CTA	Pro to Leu
amily C	Family C HNPCC Kindred	Germline	265-314	793 to 942 Deletion	Inframe Deletion
Family 8	HNPCC Kindred	Germline	406	CGA to TGA	Arg to Stop
CX 10	RER+ Tumor	Germline	639	CAT to TAT	His to Tyr
		Somatic	663	ATG to TGTG	Frameshin

Provided with the cDNA sequence and the amino acid of hMSH2 protein, one of ordinary skill in the art can readily produce hMSH2 protein, isolated and purified from other human proteins. For example, recombinant cells or organisms can be used to produce the protein in bacteria, yeast, or other convenient cell system. The isolated and purified protein can be used in screening for new therapeutic agents, for example, in in vitro assays of DNA mismatch repair. The protein can also be used to raise antibodies against hMSH2. Therapeutic administration of the protein is also contemplated.

Transgenic animals are also contemplated by the present invention. These animals would have inserted in their germline hMSH2 alleles which are associated with HNPCC or sporadic tumors. Such animals could provide model systems for testing drugs and other therapeutic agents to prevent or retard the development of tumors. Also contemplated are genetically engineered animals which contain one or more mutations in their own MSH2 genes. The mutations will be engineered to correspond to mutations found in hMSH2 alleles which are found in HNPCC-affected individuals or in other human RER+ tumors. Animals with both native MSH2 alleles inactivated and containing a human wild-type or mutant hMSH2 allele are particularly desirable.

WO 95/15381

- 13 -

Examples

Example 1

Somatic Cell Hybrids

A panel of human-hamster, human-mouse, and human-rat hybrid cell lines was developed to facilitate HNPCC mapping. Hybrids containing only portions of chromosome 2 were obtained by microcell-mediated chromosome transfer or by standard cell fusions following X-irradiation of the chromosome 2 donor. Additionally, two hybrids were used which contained a (X;2)(q28;p21) translocation derived from human fibroblasts. In previous studies, the HNPCC locus was mapped to the 25 cM region surrounding marker 123 and bordered by markers 119 and 136 (Peltomaki et al., 1993a). Thirty-eight hybrids were screened with these three chromosome 2p markers. Eight of the hybrids proved useful for mapping the relevant portion of chromosome 2p. For example, hybrids L1 and L2 contained the distal half of the region, including marker 123, while hybrid y3 contained the half proximal to marker 123 (Figure 1).

Methods

Methods for the derivation of microcell-mediated chromosome 2 hybrids have been described previously (Chen et al., 1992; Spurr et al., 1993). Some hybrids were generated following fusion of X-irradiated donor cells containing human chromosome 2 to CHO cells (Chen et al., 1994). Mouse hybrids were derived by fusing HPRT deficient L cells (A9) with human fibroblasts (GM7503) containing a t(X;2)(q28,p21) translocation and selecting in media containing HAT.

Example 2

Polymorphic Markers

To map more finely the HNPCC locus, additional polymorphic markers were obtained in three ways. First, a genomic clone containing 85 kb surrounding the 123 marker was used for fluorescence in situ hybridization (FISH) to localize it to chromosomal band 2p16.3 (Figure 2A,B). The 2p16 band region was then microdissected, and the sequences within this band were amplified using the polymerase chain reaction and subcloned into plasmid vectors (see Experimental

Procedures). The accuracy of the microdissection was confirmed using dual-color FISH by simultaneously hybridizing to microdissected material and a genomic clone containing marker 123 (Figure 2C,D). The subclones were screened by hybridization to a (CA)₁₅ probe, and hybridizing clones identified and sequenced. These sequences were then used to design oligonucleotide primers for PCR analysis of genomic DNA. Nineteen (CA), repeat markers were identified in this way. Of these, four were highly polymorphic and mapped to the region between markers 119 and 136, as assessed by the somatic cell hybrid panel exhibited in Figure 1. Second, eight additional (CA), markers, cloned randomly from human genomic DNA using a poly (CA) probe, were found to lie between markers 119 and 136 by linkage analysis in CEPH pedigrees. Five of these were particularly informative and were used in our subsequent studies. Finally, one additional marker was identified by screening subclones of a genomic P1 clone containing marker 123 with a (CA), probe. Through these analyses, thirteen new polymorphic markers were identified in the 25 cM interval between markers 119 and 136, resulting in an average marker spacing of -2 cM (Table II). These markers were mapped with respect to one another by linkage in CEPH and HNPCC pedigrees as well as by analysis of somatic cell hybrids. These two mapping techniques provided consistent and complementary information. For example, the relative positions of CA16 and CA18 could not be distinguished through linkage analysis but could be determined with the somatic cell hybrids L1, L2, and Y3. Conversely, the relative position of the ze3 and yh5 markers could not be determined through somatic cell hybrid mapping, but could be discerned by linkage analysis.

=	-	_
Ł	1	1
_	_	Ī
(1)
	1	
ŀ	_	_

	P1	CLONES							210, 211									387,388, 389
	YAC	CLONES		11E1	4F4, 1E1	4F4, 1E1, 9H6, 4A10	7F10, 4E2, 5A11	688	3D11, 8C7	3D11, 8C7	8E5	8E5	284					
TABLE II		HETEROZYGOSITY							0.78						0.79	0.73	0.73	0.78
		1 00		5.5	15.4	i	4.7	60	8.9		17.1	i	3.0	17.6		9.6		
		W	i	6.1	6.4	0.0	2.1	1.7	2.4	0.0	4.3	0.0	1.1	2.7	0.0	2.6	7	-
		DERIVATION	 	-	-	۰	≆	Œ	-	۵.	×	Σ	}-	-	- -	-	-	-
		MARKER	177 (AFM267zc9)	118 (AFM077yb7)	yh5 (AFM337yh5)	ze3 (AFM200ze3)	CAS (CAS)	CA7 (CA7)	123 (AFM083xh3)	CA2 (CA2)	CA18 (CA18)	CA16 (CA16)	yb9 (AFM320yb9)	x(5 (AFM310x(5)	xf1 (AFM348If1)	147 (AFM199vb8)	136 (AFM172xe7)	134 (AFM 168xg11)
							S	UB	ST	ITU	ITE	Sł	ΙĖΕ	ET (RL	ΙLΕ	28	5 <u>)</u>

Methods

All markers were obtained by screening human genomic libraries with radiolabelled (CA), probes (Weber and May, 1989). The "T" markers (see Table 1) were generated from a library made from total human genomic DNA, as described in Weissenbach et al., 1992. The "M" markers were made from libraries generated from microdissected chromosome 2p16, as described below. The CA2 marker was generated from a library made from P1 clone 210 digested to completion with Sau3 and cloned into the XhoI site of lambda YES (Elledge et al., 1991). The sequences of the clones obtained from these libraries were determined, and primers surrounding the CA repeats chosen. Only primers giving robust amplification and high heterozygosity were used for detailed analysis of HNPCC kindreds. All markers used in this study were shown to be derived from chromosome 2p by both linkage analysis in the CEPH pedigrees and evaluation in the somatic cell hybrid panel shown in Figure 1. The sequences of the primers and other details specific for each marker have been deposited with the Genome Data Base. Linkage analyses to obtain the map of the marker loci in CEPH families 1331, 1332, 1347, 1362, 1413, 1416, 884, and 102 (Weissenbach et al., 1992) were performed using the CLINK program of the LINKAGE program package (Lathrop et al., 1984) with the no sex difference option and 11-point computations. The odds for the best locus order supported by the data were evaluated against pairwise inversions of the loci.

Example 3

Genomic Clones

Many of the polymorphic markers shown in Table 1 were used to derive genomic clones containing 2p16 sequences. Genomic clones were obtained by PCR screening of human P1 and YAC libraries with these polymorphic markers, with ten additional sequence tagged sites (STS) derived from chromosome 2p16 microdissection, or with YAC junctions. Twenty-three P1 clones, each containing 85-95 kb, were obtained, as well as 35 YAC clones, containing 300 to 1800 kb. The YAC clones in some cases confirmed the linkage and somatic cell hybrid

maps. For example, markers ze3 and yh5 were both found in YAC's 4F4 and 1E1 while CA16 and CA18 were both found in YAC 8E5, documenting their proximity. The highest density of genomic clones (28 YAC and 17 P1 clones) was obtained between markers yb9 and yh5 (Table 1), which became the region most likely to contain the HNPCC gene during the course of these studies. The region between yh5 and yb9 was predicted to contain ~ 9 Mb (assuming 1 Mb/per cM). Based on the sizes of the YAC clones, and taking into account their chimerism, we estimated that they contained over 70% of the sequences between yh5 and yb9.

Methods

The markers described in Table 1 were used to screen YAC or P1 libraries by PCR. The CEPH A library was obtained from Research Genetics, Inc. and consisted of 21,000 YAC clones, arrayed in a format allowing facile screening and unambiguous identification of positive clones. The sizes of ten of the YAC clones containing markers were determined by transverse alternating pulse-field gel electrophoresis using a GeneLine II apparatus from Beckman and found to average 0.7 Mb (range 0.2-1.8 Mb). In some cases, inverse PCR was used to determine the YAC junctions (Joslyn et al., 1991), and the derived sequence used for "chromosome walking" with the YAC or P1 libraries. The junctions were also used to design primers to test whether the ends of the YAC clones could be localized to chromosome 2p16 (and therefore presumably be non-chimeric). Three of four YAC clones which were tested in this way had both ends within the expected region of chromosome 2, as judged by analysis with the somatic cell hybrid panel. The human genomic P1 library was also screened by PCR (Genome Systems, Inc.). P1 clones M1015 and M1016, containing the hMSH2 gene, were used to determine intron-exon borders using sequencing primers from the exons and SequiTherm™ polymerase (Epicentre Technologies).

Example 4

Analysis of HNPCC Families

The markers described in Table 1 were then used to analyze six large HNPCC kindreds previously linked to chromosome 2p (Peltomaki et al., 1993a).

Two hundred thirteen individuals, including 56 members affected with colorectal or endometrial cancer, were examined. Four of the kindreds were from the United States, one from Newfoundland and one from New Zealand. To increase the number of affected individuals that could be examined, we obtained formalin-fixed, paraffin-embedded sections of normal tissues from deceased individuals and purified DNA from them (Goelz et al., 1985a). A single allele of each of the thirteen markers was found to segregate with disease in each of the six families (i.e., the allele was found in over 50% of affected individuals). No allele of any marker was shared among the affected members of more than three kindreds.

Fourteen of the affected members contained only a subset of the expected alleles and therefore had undergone recombination between markers 119 and 136. Eleven of these individuals appeared to have simple, single recombination events. The most informative of these was in individual 148 from the J kindred and individual 44 from kindred 621 (Figure 3). Individual 621-44 apparently retained the disease linked allele at markers distal to CA5, while demonstrating multiple recombinants at more proximal loci, thus placing the CA5 marker at the proximal border of the HNPCC locus. Individual J-148 apparently retained the disease-linked allele at all markers proximal to yh5, while exhibiting recombinants at yh5 and 119, thus placing the distal border at yh5. Assuming that the same gene was involved in both the J and 621 kindreds, the HNPCC gene was predicted to reside between markers CA5 and yh5, an area spanning approximately 2 cM (Table 1).

The DNA of three affected individuals (C-202, 4-156, 4-92) appeared to have undergone two recombinations in the area. There was probably one recombinant per generation, and this could be demonstrated in C-202 by analysis of DNA from his parents; in the other cases, parental DNA was not available. All three individuals retained disease-linked alleles at CA5 and ze3 but not at more proximal and distal loci (Figure 3). Combined with the data from the patients with single recombinations, the double recombinants suggested that the HNPCC gene resided between CA5 and ze3, a distance spanning less than 2 cM.

SUBSTITUTE SHEET (RULE 26)

To determine the physical distances separating CA5, ze3, and yh5, metaphase and interphase FISH analysis was carried out. Dual-color FISH with P1 clones containing these markers was performed with P1 clones 820 and 838 (containing the markers CA5 and yh5, respectively) labeled with biotin and detected with fluorescein-labelled avidin, and clone 836 (containing marker ze3) labelled with Spectrum-Orange (Meltzer et al., 1992). The hybridization signals of these markers appeared coincident on metaphase chromosomes, confirming that they resided within an interval of <1.0 Mb. When FISH was performed on interphase nuclei, the relative positions of the three markers could be determined and the distances between them estimated (Trask et al., 1989). The results confirmed that the orientation of the markers was telomere-yh5-ze3-CA5centromere (data not shown). Direct measurement of the distance between yh5 and ze3 was estimated at < 0.3 Mb, consistent with the presence of both of these markers on YAC clones 4E4 and 1E1. Measurements of 48 interphase chromosomes provided an estimate of the distance between ze3 and CA5 at < 0.8Mb, independently confirming the linkage data.

Methods

G-banded metaphase chromosomes were microdissected with glass microneedles and amplified by PCR as previously described (Guan et al., 1993 Kao and Yu, 1991). For dual-color FISH, the PCR product was fluorochrome labelled (Spectrum-Orange, Imagenetics, Naperville, IL) or biotinylated in a secondary PCR reaction. P1 clones were labelled by nick-translation or by degenerate oligonucleotide primers (Guan et al., 1993). FISH was carried out as previously described (Guan et al., 1993) and visualized with a Zeiss Axiophot equipped with a dual-bandpass filter. For analysis of interphase FISH patterns, the distance between hybridization signals was measured in a minimum of 24 nuclei (Trask et al., 1989).

Example 5

Candidate Genes

On the basis of the mapping results described above, we could determine whether a given gene was a candidate for HNPCC by determining its position relative to the CA5-ze3 domain. The first gene considered was a human homolog of the Drosophila SOS gene (reviewed by Egan and Weinberg, 1993). This gene transmits signals from membrane bound receptors to the ras pathway in diverse eukaryotes. It was considered a candidate because another ras-interacting gene, NF1, causes a cancer predisposition syndrome (Viskochil et al., 1990; Wallace et al., 1990), and SOS has been localized to chromosome 2p16-21 by in situ hybridization (Webb et al., 1993). Using PCR to amplify SOS sequences from the hybrid panel, however, SOS was found to be distal to the CA5-yh5 domain (present in hybrid Z19 but not Z29).

We next examined the interferon-inducible RNA activated protein kinase gene PKR. This gene has been shown to have tumor suppressor ability (reviewed by Lengyel, 1993) and to map close to 2p16 (Hanash et al., 1993). We could not initially exclude PKR from the HNPCC domain, and therefore determined the sequence of its coding region in two individuals from HNPCC kindred C. Reverse transcriptase was used to generate cDNA from lymphoblastoid derived RNA of these two individuals, and PCR performed with primers specific for PKR. The PKR products were sequenced, and no deviations from the published sequence was identified within the coding region (Meurs et al., 1990). Subsequent studies showed that the PKR gene was distal to the yh5 marker, and thus could be excluded as a basis for HNPCC.

We then considered human homologs of the MutL and MutS mismatch repair genes previously shown to produce microsatellite instability in bacteria and yeast when disrupted (Levinson and Gutman, 1987; Strand et al., 1993). A human homolog of the yeast MutL-related gene PMS1 (Kramer et al., 1989) does not appear to reside on chromosome 2p (M. Liskay, personal communication). To identify homologs of MutS, we used degenerate oligonucleotide primers to PCR-

amplify cDNA from colon cancer cell lines. The same primers had been previously used to identify the yeast MSH2 gene on the basis of its MutS homology (Reenan and Kolodner, 1992). Under non-stringent conditions of PCR, a fragment of the expected size was obtained and these fragments were cloned into plasmid vectors. Most of the clones contained ribosomal RNA genes, representing abundant transcripts with weak homology to the degenerate primers. A subset of the clones, however, contained sequences similar to that of the yeast MSH2 gene, and one such clone, pNP-23, was evaluated further. The human gene from which this clone is derived is hereafter referred to as hMSH2.

The insert from clone pNP-23 was used as a probe in Southern blots of somatic cell hybrid DNA. This insert hybridized to one or two fragments in human genomic DNA digested with PstI or EcoRI, respectively, and these fragments were present in hybrid Z30, containing most of human chromosome 2p. Analysis of other hybrids showed that the fragment was present in hybrids Z11, Z29, L1 and L2, but not Z12, Y3 or Z19, thereby localizing the human MSH2 (hMSH2) gene to a region bordered by markers CA18 and 119 (examples in Figure 4). The YAC clones listed in Table 1 were then analyzed, and EcoRI and PstI fragments of the expected size identified in YAC 5A11, derived from screening the YAC library with the CA5 marker (Figure 4).

To confirm the Southern blots, we designed non-degenerate primers on the basis of the sequence of pNP-23. Several sets of primers were tested so that genomic DNA could be used as a template for PCR; an intervening intron prevented the original primers from being used effectively with templates other than cDNA. PCR with these primers was perfectly consistent with the Southern blot results. The expected 101 bp fragment was present in hybrids Z4, Z11, Z29, L1, and L2, and in YAC 5A11, but not in other hybrids or YAC clones (not shown).

The localization of hMSH2 sequences to YAC 5A11 demonstrated the proximity of these sequences to marker CA5. To determine the distance and relative orientation of hMSH2 with respect to CA5, we performed interphase FISH

analysis. P1 clones containing CA5, ze3 and hMSH2 sequences (clones 820, 836, and M1015, respectively) were simultaneously hybridized to interphase nuclei following fluorescein and Spectrum Orange labelling (Meltzer et al., 1992). The results demonstrated that MSH2 resides within the HNPCC locus defined by linkage analysis to lie between CA5 and ze3, and less than 0.3 Mb from marker CA5 (Figure 2E).

cDNA libraries generated from human colon cancer cells or from human fetal brain tissues were then screened with the insert of pNP-23 to obtain additional sequences from this gene. Seventy-five cDNA clones were initially identified and partially sequenced. PCR products representing the ends of the cDNA sequence contig were then used as probes to re-screen the cDNA libraries. This cDNA "walk" was repeated again with the new contig ends. Altogether, 147 cDNA clones were identified. The composite sequence derived from these clones is shown in Figure 5. An open reading frame (ORF) began 69 nt downstream of the 5' end of the cDNA contig, and continued for 2802 bp. The methionine initiating this ORF was in a sequence context compatible with efficient translation (Kozak, 1986) and was preceded by in-frame termination codons. RNA from placenta and brain were used in a PCR-based procedure (RACE, Frohman et al., 1988) to independently determine the position of the 5' end of hMSH2 transcripts. This analysis demonstrated that the 5' ends of all detectable transcripts were less than 100 bp upstream of the sequence shown in Figure 5, and were heterogeneous upstream of nt -69. The region of highest homology to the yeast MSH2 gene is shown in Figure 6. This region encompassed the helix-turn-helix domain perhaps responsible for MutS binding to DNA (Reenan and Kolodner, 1992). The yeast and human MSH2 proteins were 77% identical between codons 615 and 788. There were several other blocks of similar amino acids distributed throughout the length of these two proteins (966 and 934 amino acids in yeast and human, respectively).

Methods

cDNA generated from the RNA of colorectal cancer cells with reverse transcriptase was used as template for PCR with the degenerate primers 5'-CTG GAT CCA C(G/A/T/C) G G(G/A/T/C)C C(G/A/T/C)A A(T/C)A TG-3' and 5'-CTG GAT CC(G/A) TA(G/A) TG(G/A/T/C) GT (G/A/T/C) (G/A) C(G/A) AA-3'. These two primers were used previously to identify the yeast MSH2 gene and were based on sequences conserved among related mammalian and bacterial genes (Reenan and Kolodner, 1992). The optimal PCR conditions for detecting the human MSH2 gene consisted of 35 cycles at 95° for 30 seconds, 41° for 90 seconds, and 70° for 90 seconds, in the buffer described previously (Sidransky et al., 1991). PCR products were cloned into T-tailed vectors as described (Holton and Graham, 1991) and sequenced with modified T7 polymerase (USB). The insert from one clone (pNP-23) containing human sequences homologous to the yeast MSH2 gene was then used to screen cDNA libraries generated from RNA of SW480 colon cancer cells (Clontech) or of fetal brain (Stratagene). After two further rounds of screening, positive clones were converted into plasmids and sequenced using modified T7 polymerase (Kinzler et al., 1991). In some cases, the inserts were amplified using one hMSH2-specific primer and one vectorspecific primer, and then sequenced with SequiTherm Polymerase (Epicentre To determine the 5' end of MSH2 transcripts, RACE was Technologies). performed (Frohman et al., 1989) using brain and placenta cDNA (Clontech).

Example 6

Mutations of hMSH2

The physical mapping of hMSH2 to the HNPCC locus was intriguing but could not prove that this gene was responsible for the disease. To obtain more compelling evidence, we determined whether germ line mutations of hMSH2 were present in the two HNPCC kindreds that originally established linkage to chromosome 2 (Peltomaki et al., 1993a). Intron-exon borders within the most conserved region of hMSH2 (Figure 6) were determined by sequencing genomic PCR fragments containing adjacent exons. Genomic DNA samples from the lymphocytes of affected members of these two kindreds were then used as

templates for PCR to determine the sequence of this domain. The DNA from individual J-42, afflicted with colon and endometrial cancer at ages 42 and 44, respectively, was found to contain one allele with a C to T transition at codon 622 (CCA to CTA), resulting in a substitution of leucine for proline (Figure 7, top). Twenty additional DNA samples from unrelated individuals all encoded proline at this position. Twenty one members of the J kindred were then analyzed by direct sequencing of PCR products. All eleven affected individuals contained one allele with a C to T transition in codon 622, while all ten unaffected members contained two normal alleles, thus documenting perfect segregation with disease. Importantly, this proline was at a highly conserved position, the identical residue being found in all known MutS related genes from prokaryotes and eukaryotes (Figure 6 and Reenan and Kolodner, 1992).

No mutations of the conserved region of MSH2 were identified in kindred C, so we next examined other parts of the hMSH2 transcript. RNA was purified from lymphoblastoid cells of patient C-202, a 27 year old male with colon cancer. Reverse transcriptase coupled PCR (RT-PCR) was used to generate four hMSH2-specific products encompassing codons 89 to 934 from this RNA (see Experimental Procedures). An abnormal, smaller RT-PCR product was identified with one of the primer pairs used. Mapping and sequencing studies using various MSH2 primers showed that the abnormal product was the result of a presumptive splicing defect which removed codons 265 to 314 from the hMSH2 transcript. The abnormal transcript was found to segregate with disease in the C kindred, and was not found in twenty unrelated individuals.

We next wished to determine whether hMSH2 was altered in one of the more recently linked families (R.P. and M.N-L., unpublished data), and chose kindred 8 for detailed analysis. DNA and RNA were obtained from lymphoblastoid cells of 8-143, a 42 year old male with colon cancer. The conserved region of hMSH2 was amplified from genomic DNA using PCR and directly sequenced. A T to C substitution was noted in the polypyrimidine tract upstream of the exon beginning at codon 669 (at intron position -6). However,

this substitution was also found in two of twenty unrelated, normal individuals, and was therefore a polymorphism unrelated to the disease, with an allele frequency of 0.05. Most of the hMSH2 coding region was then amplified by RT-PCR, as described above, and no abnormal transcripts were detected. Sequencing of the PCR products, however, revealed a C to T transition at codon 406 (CGA to TGA) causing substitution of a termination codon for an arginine residue. RNA was available from the lymphocytes of a second affected member of kindred 8, and the same stop codon was identified. This alteration was not found in twenty other, unrelated individuals.

Finally, we wished to determine whether mutations of this gene occurred in RER+ tumors from patients without evident family histories of cancer. The conserved region of MSH2 was studied in four colorectal tumor cell lines from such patients using genomic DNA as templates for PCR. One tumor (from patient Cx10) was found to contain two hMSH2 alterations. The first was a C to T transition in codon 639 (CAT to TAT), resulting in a substitution of tyrosine for histidine. This change was not found in any of twenty samples from unrelated individuals, but was present in the DNA from normal colon of this patient, and was therefore likely to represent a germ line change (Figure 7, middle). Like the missense mutation in the J kindred, the Cx10 alteration was at a position perfectly conserved in all MutS homologs (Figure 6 and Reenan and Kolodner, 1992). The second alteration in the tumor from Cx10 was a substitution of a GT dinucleotide for an A in codon 663 (ATG to TGTG). The resultant one bp insertion was predicted to cause a frameshift, producing a termination codon 36 nt downstream. This mutation was demonstrated in both RNA and DNA purified from the Cx10 tumor, but was not present in the patient's normal colon, so represented a somatic mutation (Figure 7, bottom). The PCR products from Cx10 were cloned and sequenced, and the insertion mutation at codon 663 and the transition at codon 639 were shown to reside on different alleles.

Methods

To detect mutations, PCR products were generated from cDNA and human genomic DNA templates, then sequenced directly using SequiTherm™. In some cases, the PCR products were cloned into T-tailed vectors for sequencing to confirm the direct sequencing data. The primers used to amplify the conserved region of MSH2 from genomic DNA were 5'-CCA CAA TGG ACA CTT CTG C-3' and 5'-CAC CTG TTC CAT ATG TAC G-3', resulting in a 1.4 kb fragment containing hMSH2 codons 614 to 705, and primers 5'-AAA ATG GGT TGC AAA CAT GC-3' and 5'-GTG ATA GTA CTC ATG GCC C-3', resulting in a 2.0 kb fragment containing MSH2 cDNA codons 683 to 783. Primers for RT-PCR were 5'-AGA TCT TCT TCT GGT TCG TC-3' and 5'-GCC AAC AAT AAT TTC TGG TG-3' for codons 89 to 433, 5'-TGG ATA AGA ACA GAA TAG AGG-3' and 5'-CCA CAA TGG ACA CTT CTG C-3' for codons 350-705, 5'-CAC CTG TTC CAT ATG TAC G-3' and 5'-AAA ATG GGT TGC AAA CAT GC-3' for codons 614 to 783, and 5'-GTG ATA GTA CTC ATG GCC C-3' and 5'-GAC AAT AGC TTA TCA ATA TTA CC-3' for codons 683-949.

Discussion

Three major conclusions can be drawn from the examples described here. First, physical mapping and linkage analysis localized the HNPCC locus on chromosome 2 to an 0.8 Mb segment bordered by markers CA5 and ze3. Second, a new human homolog of the yeast MSH2 gene was identified, and this gene shown to lie in the same 0.8 Mb interval. Third, alterations of the hMSH2 gene occurred in the germ line of patients with RER⁺ tumors, with or without classical HNPCC, and additional somatic alterations of this gene occurred in tumors (Summarized in Table I). The alterations were at highly conserved regions or significantly altered the expected gene product and thus represent mutations with important functional effects. These results indicate that mutations of hMSH2 are responsible for HNPCC and the RER⁺ positive phenotype found in tumors.

SUBSTITUTE SHEET (RULE 26)

These data have substantial implications for understanding the neoplastic disease observed in HNPCC. In particular, they suggest that the microsatellite alterations previously observed in tumors from these patients are not epiphenomena, but are intrinsically related to pathogenesis. Additionally, the mutations observed in yeast and bacteria with defective MutS-related genes are not confined to insertions and deletions at simple repeated sequences, though these sequences provide convenient tools for analysis (Modrich, 1991). Similarly, one would expect that many mutations, in addition to microsatellite insertions or deletions, would be found in HNPCC tumors. This could lead to the multiple, sequential mutations in oncogenes and tumor suppressor genes which have been shown to drive colorectal tumorigenesis (Fearon and Vogelstein, 1990). Thus, the molecular pathogenesis of HNPCC tumors is likely to be similar to that occurring in non-HNPCC cases, though accelerated by the increased rate of mutation associated with mismatch repair defects. Accordingly, colon tumors from HNPCC patients have been shown to contain mutations of APC, p53 and RAS at frequencies similar to those found in sporadic colorectal cancers (Aaltonen et al., 1993).

Colorectal tumors from HNPCC patients are distinguished by their relatively normal cytogenetic composition (Kouri et al., 1990), and sporadic, RER+ tumors have been demonstrated to have substantially fewer chromosome losses than those occurring in RER cases (Thibodeau et al., 1993; Aaltonen et al., 1993). These data suggest that genetic heterogeneity is critical for colorectal cancer development, but can be generated in two different ways (Thibodeau et al., 1993). Most commonly, it develops through gross alterations resulting in aneuploidy, as suggested nearly eighty years ago (Boveri, 1914). In HNPCC-derived tumors and RER+ sporadic tumors, the diversity is presumably more subtle, consisting of multiple small sequence changes distributed throughout the genome. The latter mechanism of generating diversity may be less dangerous to the host, as HNPCC patients, as well as patients with RER+ sporadic tumors, appear to have a better prognosis than would be expected from histopathologic

analysis of their tumors (Ionov et al.,; 1993; Thibodeau et al., 1993; Lothe et al., 1993; Lynch et al., 1993).

References

Aaltonen, L.A., Peltomaki, P., Leach, F.S., Sistonen, P., Pylkkanen, L., Mecklin, J-P., Jarvinen, H., Powell, S.M., Jen, J., Hamilton, S.R., Petersen, G.M., Kinzler, K.W., Vogelstein, B., and de la Chapelle, A. (1993). Clues to the pathogenesis of familial colorectal cancer. Science 260, 812-816.

Baylin, S.B., Makos, M., Wu, J., C-Y, R-W., de Bustros, A., Vertino, P., and Nelkin, B.D. (1991). Abnormal patterns of DNA methylation in human neoplasia: Potential consequences for tumor progression. Cancer Cells 3, 383, 390.

Boveri, T. (1914). Zur Frage der Entstehung maligner tumoren. Gustave Fischer Verlag, Jena, Vol. 1.

Chen, D.J., Park, M.S., Campbell, E., Oshimura, M., Liu, P., Zhao, Y., White, B.F., and Siciliano, M.J. (1992). Assignment of a human DNA double-strand break repair gene (XRCC5) to chromosome 2. Genomics 13, 1088-1094.

Chen, D.J., Marrone, B., Nguyen, T., Stackhouse, M., Zhao, Y., and Sicilano, J.J., (1994). Regional assignment of a human radiation repair gene (XRCC5) to 2q35 by X-ray hybrid mapping. Genomics, in press.

Egan, S.E., and Weinberg, R.A. (1993). The pathway to signal achievement. Nature 365, 781-783.

Elledge, S.J., Mulligan, J.T., Ramer, S.W., Spottswood, M., and Davis R.W. (1991). Lambda YES: a multifunctional cDNA expression vector for the isolation

SUBSTITUTE SHEET (RULE 26)

of genes by complementation of yeast and Escherichia coli mutations. Proc. Natl. Acad. Sci. USA 88, 1731-1735.

Fang, W.-h. and P. Modrich (1993). Human strand-specific mismatch repair occurs by a bidirectional mechanism similar to that of the bacterial reaction. J. Biol. Chem. 268, 11838-11844.

Fearon, E.R., and Vogelstein, B. (1990). A genetic model for colorectal tumorigenesis. Cell 61, 759-767.

Frohman, M.A, Dush, M.K., and Martin, G.R. (1988). Rapid production of full-length cDNAs from rare transcripts: Amplification using a single gene-specific oligonucleotide primer. Proc. Natl. Acad. Sci. USA 85, 8998-9002.

Goelz, S.E., Hamilton, S.R., and Vogelstein, B. Purification of DNA from Formaldehyde Fixed and Paraffin Embedded Human Tissue. (1985a). Biochem. Biophys. Res. Commun. 130, 118-126.

Goelz, S.E., Vogelstein, B., Hamilton, S.R., and Feinberg, A.P. Hypomethylation of DNA from Benign and Malignant Human Colon Neoplasms. (1985b). Science 228, 187-190.

Guan, X.-Y., Trent, J.M., and Meltzer, P.S. (1993). Generation of band-specific painting probes from a single microdissected chromosome. Human Molecular Genetics 2, 1117-1121.

Han, H-J., Yanagisawa, A., Kato, Y., Park, J-G., and Nakamura, Y. (1993). Genetic instability in pancreatic cancer and poorly differentiated type of gastric cancer. Cancer Research 53, 5087-5089.

Hanash, S.M., Beretta, L., Barcroft, C.L., Sheldon, S., Glover, T.W., Ungar, D., and Sonenberg, N. (1993). Mapping of the gene for interferon-inducible dsRNA-dependent protein kinase to chromosome region 2p21-22: A site of rearrangements in myeloproliferative disorders. Genes, Chromosomes & Cancer 8, 34-37.

Holmes, J., S. Clark and P. Modrich (1990). Strand-specific mimsatch correction in nuclear extracts of human and *Drosophila melanogaster* cell lines. Proc. Natl. Acad. Sci. U.S.A. 87, 5837-5841.

Holton, T.A., and Graham, M.W. (1991). A simple and efficient method for direct cloning of PCR products using ddT-tailed vectors. Nucleic Acids Research 19, 1156.

Ionov, Y.M., Peinado, A., Malkhosyan, S., Shibata, D., and Perucho M. (1993). Ubiquitous somatic mutations in simple repeated sequences reveal a new mechanism for colonic carcinogenesis. Nature 363, 558-561.

Jass, J.R., Stewart, S.M. (1992). Evolution of hereditary non-polyposis colorectal cancer. Gut 33, 783-786.

Joslyn, G., Carlson, M., Thilveris, A., Albertsen, H., Gelbert, L., Samowitz, W., Groden, J., Stevens, J., Spirio, L., Robertson, M., Sargeant, L., Krapcho, K., Wolff, E., Burt, R., Hughes, J.P., Warrington, J., McPherson, J., Wasmuth, J., Le Paslier, D., Abderrahim, H., Cohen, D., Leppert, M., and White, R. (1991). Identification of deletion mutations and three new genes at the familial polyposis locus. Cell 66, 601-613.

Kao, F-T., and Yu, J-W. (1991). Chromosome microdissection and cloning in human genome and genetic disease analysis. Proc. Natl. Acad. Sci. USA 88, 1844-1848.

Kinzler, K.W., Nilbert, M.C., Su, L.-K., Vogelstein, B., Bryan, T.M., Levy, D.B., Smith, K.J., Preisinger, A.C., Hedge, P., McKechnie, D., Finniear, R., Markham, A., Groffen, J., Boguski, M.S., Altschul, S.F., Horii, A., Ando, H., Miyoshi, Y., Miki, Y., Nishisho, I., Nakamura, Y. Identification of FAP Locus Genes from Chromosome 5q21. (1991). Science 253, 661-665.

Kouri, M., Laasonen, A., Mecklin, J.P., Jarvinen, H., Franssila, K., Pyrhonen, S. (1990). Diploid predominance in hereditary nonpolyposis colorectal carcinoma evaluated by flow cytometry. Cancer 65, 1825-1829.

Kozak, M. (1986). Point mutations define a sequence flanking the AUG initiator codon that modulates translation by Eukaryotic Ribosomes. Cell 44, 283-292.

Kramer, W., Kramer, B., Williamson, M.S., Fogel, S. (1989). Cloning and nucleotide sequence of DNA mismatch repair gene PMS1 from Saccharomyces cerevisiae: homology of PMS1 to procaryotic MutL and HexB. Journal of Bacteriology 171, 5339-5346.

Knudson, A.G. (1993). All in the (cancer) family. Nature Genetics 5, 103-104.

Kunkel, T.A. (1993). Slippery DNA and diseases. Nature 365, 207-208.

Lathrop, G.M., Lalouel, J.M., Julier, C., and Ott, J. (1984). Strategies for multilocus linkage analysis in humans. Proc. Natl. Acad. Sci. USA 81, 3443-3446.

SUBSTITUTE SHEET (RULE 26)

Lengyel, P. (1993). Tumor-suppressor genes: News about the interferon connection. Proc. Natl. Acad. Sci. USA 90, 5893-5895.

Levinson, G., and Gutman, G.A. (1987). High frequencies of short frameshifts in poly-CA/TG tandem repeats borne by bacteriophage M13 in *Escherichia coli* K-12. Nucleic Acids Research 15, 5323-5338.

Lindblom, A., Tannergard, P., Werelius, B., and Nordenskjold, M. (1993). Genetic mapping of a second locus predisposing to hereditary non-polyposis colon cancer. Nature Genetics 5, 279-282.

Lothe, R.A., Peltomaki, P., Meling, G-I., Aaltonen, L.A., Nystrom-Lahti, M., Pylkkanen, L., Heimdal, K., Andersen, T.I., Moller, P., Rognum, T.O., Fossa, S.D., Haldorsen, T., Langmark, F., Bragger, A., de la Chapelle, A., and Barresea, A-L. (1993). Genomic instability in colorectal caner: Relationship to clinicopathological variables and family history. Cancer Research, in press.

Lynch, H.T., Omaha, M.W., Shaw, M.D. (1966). Hereditary factors in cancer. Arch Intern Med 117, 206-212.

Lynch, H.T., Smyrk, T.C., Watson, P., Lanspa, S.J., Lynch, J.F., Lynch, P.M., Cavalieri, R.J., and Boland, C.R. (1993). Genetics, natural history, tumor spectrum, and pathology of hereditary nonpolyposis colorectal cancer: An updated review. Gastroenterology 104, 1535-1549.

Meltzer, P.S., Guan, X-Y., Burgess, A., and Trent, J.M. (1992). Rapid generation of region specific probes by chromosome microdissection and their application. Nature Genetics 1, 24-28.

Meurs, E., Chong, K., Galabru, J., Thomas, N.S.B., Kerr, I.M., Williams, B.R.G., and Hovanessian, A.G. (1990). Molecular cloning and characterization of the human double-stranded RNA-activated protein kinase induced by interferon. Cell 62, 379-390.

Modrich, P. (1991). Mechanisms and biological effects of mismatch repair. Ann. Rev. Genet. 25, 229-253.

Peinado, M.A., Malkhosyan, S., Velazquez, A., and Perucho, M. (1992). Isolation and characterization of allelic losses and gains in colorectal tumors by arbitrarily primed polymerase chain reaction. Proc. Natl. Acad. Sci. USA 89, 10065-10069.

Peltomaki, P., Aaltonen, L.A., Sistonen, P., Pylkkanen, L., Mecklin, J-P., Jarvinen, H., Green, J.S., Jass, J.R., Weber, J.L., Leach, F.S., Petersen, G.M., Hamilton, S.R., de la Chapelle, A., and Vogelstein, B. (1993a). Genetic mapping of a locus predisposing to human colorectal cancer. Science 260, 810-812.

Pukkila, P.J., Petseon, J., Herman, G., Modrich, P., and Meselson, M. (1983). Effects of high levels of DNA adenine methylation on methyl-directed mismatch repair in *Escherichia coli*. Genetics 104, 571-582.

Reenan, R.A., Kolodner, R.D. (1992). Isolation and characterization of two Saccharomyces cerevisiae genes encoding homologs of the bacterial HexA and MutS mismatch repair proteins. Genetics 132, 963-973.

Risinger, J.I., Berchuck, A., Kohler, M.F., Watson, P., Lynch, H.T., and Boyd, J. (1993). Genetic instability of microsatellites in endometrial carcinoma. Cancer Research 53, 5100-5103.

SUBSTITUTE SHEET (RULE 26)

WO 95/15381

Sidransky, D., von Eschenbach, A., Tsai, Y.C., Jones, P., Summerhayes, I., Marshall, F., Paul, M., Green, P., Hamilton, S.R., Frost, P., Vogelstein, B. (1991). Identification of p53 Gene Mutations in Bladder Cancers and Urine Samples. Science 252, 706-709.

Spurr, N.K., Cox, S., Naylor, S. (1993). Report and abstracts of the Second International Workshop on human chromosome 2 mapping. Cytogenetics & Cell Genetics 64, 69-92.

Stanbridge, E.J. (1990). Human tumor suppressor genes. Ann. Rev. Genes 24, 615-657.

Strand, M., Prolla, T.A., Liskay, R.M., and Petes, T.D. (1993). Destabilization of tracts of simple repetitive DNA in yeast by mutations affecting DNA mismatch repair. Nature 365, 274-276.

Thibodeau, S.N., Bren, G., and Schaid, D. (1993). Microsatellite instability in cancer of the proximal colon. Science 260, 816-819.

Thomas, D.C., J.D. Roberts and T.A. Kunkel (1991). Heteroduplex repair in extracts of human HeLa cells. J.Biol. Chem. 266, 3744-3751.

Trask, B., Pinkel, D., and van den Engh, G. (1989). The proximity of DNA sequences in interphase cell nuclei is correlated to genomic distance and permits ordering of cosmids spanning 250 kilbase pairs. Genomics 5, 710-717.

Viskochil, D., Buchberg, A.M., Xu, G., Cawthon, R., Stevens, J., Wolff, R.K., Culver, M., Carey, J.C., Copeland, N.G., Jenkins, N.A., White, R., and O'Connell, P. (1990). Deletions and a translocation interrupt a cloned gene at the neurofibromatosis type 1 locus. Cell 62, 187-192.

SUBSTITUTE SHEET (RULE 26)

Vogelstein, B., Fearon, E.R., Hamilton, S.R., Preisinger, A.C., Willard, H.F., Michelson, A.M., Riggs, A.D., and Orkin, S.H. (1987). Clonal Analysis Using Recombinant DNA Probes from the X Chromosome. Cancer Research 47, 4806-4813.

Wallace, M.R., Marchuk, D.A., Andersen, L.B., Letcher, R., Odeh, H.M., Saulino, A.M., Fountain, J.W., Brereton, A., Nicholson, J., Mitchell, A.L., Brownstein, B.H., Collins, F.S. (1990). Type 1 neurofibromatosis gene: Identification of a large transcript disrupted in three NF1 patients. Science 249, 181-186.

Warthin, A.S. (1913). Heredity with reference to carcinoma: As shown by the study of the cases examined in the pathological laboratory of the University of Michigan, 1895-1913. Arch. Int. Med 12, 546-555.

Watson, P., and Lynch, H.T. (1993). Extracolonic cancer in hereditary nonpolyposis colorectal cancer. Cancer 71, 677-685.

Webb, G.C., Jenkins, N.A., Largaespada, D.A., Copeland, N.G., Fernandez, C.S., and Bowtell, D.D.L. (1993). Mammalian homologues of the Drosophila Son of sevenless gene map to murine chromosomes 17 and 12 and to human chromosomes 2 and 14, respectively. Genomics 18, 14-19.

Weber, J.L., May P.E. (1989). Abundant class of human DNA polymorphisms which can be typed using the polymerase chain reaction. American Journal of Human Genetics 44, 338-396.

Weissenbach, J., Gyapay, G., Dib, C., Vignal A., Morissette, J., Millasseau, P., Vaysseix, G., and Lathrop, M. (1992). A second-generation linkage map of the human genome. Nature 359, 794-801.

SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT: Vogelstein, Bert Kinzler, Kenneth W. de la Chappelle, Albert
- (ii) TITLE OF INVENTION: Mutator Gene and Hereditary Non-Polyposis Colorectal Cancer
- (iii) NUMBER OF SEQUENCES: 16
- (iv) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Banner, Birch, McKie, and Beckett
 - (B) STREET: 1001 G Street, N.W.
 - (C) CITY: Washington
 - (D) STATE: D.C.
 - (E) COUNTRY: USA
 - (F) ZIP: 20001
 - (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk

 - (B) COMPUTER: IBM PC compatible
 (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 08/160295
 - (B) FILING DATE: 02-DEC-1993
 - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Kagan, Sarah A.
 - (B) REGISTRATION NUMBER: 32,141
 - (C) REFERENCE/DOCKET NUMBER: 01107.44900
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: 202.508.9100

 - (B) TELEFAX: 202.508.9299 (C) TELEX: 197430 BBMB UT
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2947 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA
 - (iii) HYPOTHETICAL: NO
 - (iv) ANTI-SENSE: NO
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo sapiens

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

GGCGGGAAAC AGCTTAGTGG	GTGTGGGGTC	GCGCATTTTC	TTCAACCAGG	AGGTGAGGAG	60
GTTTCGACAT GGCGGTGCAG	CCGAAGGAGA	CGCTGCAGTT	GGAGAGCGCG	GCCGAGGTCG	120
GCTTCGTGCG CTTCTTTCAG	GGCATGCCGG	AGAAGCCGAC	CACCACAGTG	CGCCTTTTCG	180
ACCGGGGCGA CTTCTATACG	GCGCACGGCG	AGGACGCGCT	GCTGGCCGCC	CGGGAGGTGT	240
TCAAGACCCA GGGGGTGATC	AAGTACATGG	GGCCGGCAGG	AGCAAAGAAT	CTGCAGAGTG	300
TTGTGCTTAG TAAAATGAAT	TTTGAATCTT	TTGTAAAAGA	TCTTCTTCTG	GTTCGTCAGT	360
ATAGAGTTGA AGTTTATAAG	AATAGAGCTG	GAAATAAGGC	ATCCAAGGAG	AATGATTGGT	420
ATTTGGCATA TAAGGCTTCT	CCTGGCAATC	TCTCTCAGTT	TGAAGACATT	CTCTTTGGTA	480
ACAATGATAT GTCAGCTTCC	ATTGGTGTTG	TGGGTGTTAA	AATGTCCGCA	GTTGATGGCC	540
AGAGACAGGT TGGAGTTGGG	TATGTGGATT	CCATACAGAG	GAAACTAGGA	CTGTGTGAAT	600
TCCCTGATAA TGATCAGTTC	TCCAATCTTG	AGGCTCTCCT	CATCCAGATT	GGACCAAAGG	660
AATGTGTTTT ACCCGGAGGA	GAGACTGCTG	GAGACATGGG	GAAACTGAGA	CAGATAATTC	720
AAAGAGGAGG AATTCTGATC	ACAGAAAGAA	AAAAAGCTGA	CTTTTCCACA	AAAGACATTT	780
ATCAGGACCT CAACCGGTTG	TTGAAAGGCA	AAAAGGGAGA	GCAGATGAAT	AGTGCTGTAT	840
TGCCAGAAAT GGAGAATCAG	GTTGCAGTTT	CATCACTGTC	TGCGGTAATC	AAGTTTTTAG	900
AACTCTTATC AGATGATTCC	AACTTTGGAC	AGTTTGAACT	GACTACTTTT	GACTTCAGCC	960
AGTATATGAA ATTGGATATT	GCAGCAGTCA	GAGCCCTTAA	CCTTTTTCAG	GGTTCTGTTG	1020
AAGATACCAC TGGCTCTCAG	TCTCTGGCTG	CCTTGCTGAA	TAAGTGTAAA	ACCCCTCAAG	1080
GACAAAGACT TGTTAACCAG	TGGATTAAGC	AGCCTCTCAT	GGATAAGAAC	AGAATAGAGG	1140
AGAGATTGAA TTTAGTGGAA	GCTTTTGTAG	AAGATGCAGA	ATTGAGGCAG	ACTTTACAAG	1200
AAGATTTACT TCGTCGATTC	CCAGATCTTA	ACCGACTTGC	CAAGAAGTTT	CAAAGACAAG	1260
CAGCAAACTT ACAAGATTGT	TACCGACTCT	ATCAGGGTAT	AAATCAACTA	CCTAATGTTA	1320
TACAGGCTCT GGAAAAACAT	GAAGGAAAAC	ACCAGAAATT	ATTGTTGGCA	GTTTTTGTGA	1380
CTCCTCTTAC TGATCTTCGT	TCTGACTTCT	CCAAGTTTCA	GGAAATGATA	GAAACAACTT	1440
TAGATATGGA TCAGGTGGAA	AACCATGAAT	TCCTTGTAAA	ACCTTCATTT	GATCCTAATC	1500
TCAGTGAATT AAGAGAAATA	ATGAATGACT	TGGAAAAGAA	GATGCAGTCA	ACATTAATAA	1560
GTGCAGCCAG AGATCTTGGC	TTGGACCCTG	GCAAACAGAT	TAAACTGGAT	TCCAGTGCAC	1620
AGTTTGGATA TTACTTTCGT	GTAACCTGTA	AGGAAGAAAA	AGTCCTTCGT	AACAATAAAA	1680
ACTITAGTAC TGTAGATATC	CAGAAGAATG	GTGTTAAATT	TACCAACAGC	AAATTGACTT	1740
CTTTAAATGA AGAGTATACC	AAAAATAAAA	CAGAATATGA	AGAAGCCCAG	GATGCCATTG	1800

TTAAAGAAAT	TGTCAATATT	TCTTCAGGCT	ATGTAGAACC	AATGCAGACA	CTCAATGATG	1860
TGTTAGCTCA	GCTAGATGCT	GTTGTCAGCT	TTGCTCACGT	GTCAAATGGA	GCACCTGTTC	1920
CATATGTACG	ACCAGCCATT	TTGGAGAAAG	GACAAGGAAG	AATTATATTA	AAAGCATCCA	1980
GGCATGCTTG	TGTTGAAGTT	CAAGATGAAA	TTGCATTTAT	TCCTAATGAC	GTATACTTTG	2040
AAAAAGATAA	ACAGATGTTC	CACATCATTA	CTGGCCCCAA	TATGGGAGGT	AAATCAACAT	2100
ATATTCGACA	AACTGGGGTG	ATAGTACTCA	TGGCCCAAAT	TGGGTGTTTT	GTGCCATGTG	2160
AGTCAGCAGA	AGTGTCCATT	GTGGACTGCA	TCTTAGCCCG	AGTAGGGGCT	GGTGACAGTC	2220
AATTGAAAGG	AGTCTCCACG	TTCATGGCTG	AAATGTTGGA	AACTGCTTCT	ATCCTCAGGT	2280
CTGCAACCAA	AGATTCATTA	ATAATCATAG	ATGAATTGGG	AAGAGGAACT	TCTACCTACG	2340
ATGGATTTGG	GTTAGCATGG	GCTATATCAG	AATACATTGC	AACAAAGATT	GGTGCTTTTT	2400
GCATGTTTGC	AACCCATTTT	CATGAACTTA	CTGCCTTGGC	CAATCAGATA	CCAACTGTTA	2460
ATAATCTACA	TGTCACAGCA	CTCACCACTG	AAGAGACCTT	AACTATGCTT	TATCAGGTGA	2520
AGAAAGGTGT	CTGTGATCAA	AGTTTTGGGA	TTCATGTTGC	AGAGCTTGCT	AATTTCCCTA	2580
AGCATGTAAT	AGAGTGTGCT	AAACAGAAAG	CCCTGGAACT	TGAGGAGTTT	CAGTATATTG	2640
GAGAATCGCA	AGGATATGAT	ATCATGGAAC	CAGCAGCAAA	GAAGTGCTAT	CTGGAAAGAG	2700
AGCAAGGTGA	AAAAATTATT	CAGGAGTTCC	TGTCCAAGGT	GAAACAAATG	CCCTTTACTG	2760
AAATGTCAGA	AGAAAACATC	ACAATAAAGT	TAAAACAGCT	AAAAGCTGAA	GTAATAGCAA	2820
AGAATAATAG	CTTTGTAAAT	GAAATCATTT	CACGAATAAA	AGTTACTACG	TGAAAAATCC	2880
CAGTAATGGA	ATGAAGGTAA	TATTGATAAG	CTATTGTCTG	TAATAGTTTT	ATATTGTTTT	2940
ATATTAA						2947

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 934 amino acids
 (B) TYPE: amino acid
 (C) STRANDEDNESS: single

 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (iii) HYPOTHETICAL: YES
 - (iv) ANTI-SENSE: NO
 - (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Homo sapiens
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:
 - Met Ala Val Gln Pro Lys Glu Thr Leu Gln Leu Glu Ser Ala Ala Glu 10

Val Gly Phe Val Arg Phe Phe Gln Gly Met Pro Glu Lys Pro Thr Thr Thr Val Arg Leu Phe Asp Arg Gly Asp Phe Tyr Thr Ala His Gly Glu Asp Ala Leu Leu Ala Ala Arg Glu Val Phe Lys Thr Gln Gly Val Ile 50 60 Lys Tyr Met Gly Pro Ala Gly Ala Lys Asn Leu Gln Ser Val Val Leu Ser Lys Met Asn Phe Glu Ser Phe Val Lys Asp Leu Leu Leu Val Arg Gln Tyr Arg Val Glu Val Tyr Lys Asn Arg Ala Gly Asn Lys Ala Ser 105 Lys Glu Asn Asp Trp Tyr Leu Ala Tyr Lys Ala Ser Pro Gly Asn Leu Ser Gln Phe Glu Asp Ile Leu Phe Gly Asn Asn Asp Met Ser Ala Ser 135 Ile Gly Val Val Gly Val Lys Met Ser Ala Val Asp Gly Gln Arg Gln Val Gly Val Gly Tyr Val Asp Ser Ile Gln Arg Lys Leu Gly Leu Cys Glu Phe Pro Asp Asn Asp Gln Phe Ser Asn Leu Glu Ala Leu Leu Ile 185 Gln Ile Gly Pro Lys Glu Cys Val Leu Pro Gly Gly Glu Thr Ala Gly 200 Asp Met Gly Lys Leu Arg Gln Ile Ile Gln Arg Gly Gly Ile Leu Ile Thr Glu Arg Lys Lys Ala Asp Phe Ser Thr Lys Asp Ile Tyr Gln Asp Leu Asn Arg Leu Leu Lys Gly Lys Lys Gly Glu Gln Met Asn Ser Ala Val Leu Pro Glu Met Glu Asn Gln Val Ala Val Ser Ser Leu Ser Ala 265 Val Ile Lys Phe Leu Glu Leu Leu Ser Asp Asp Ser Asn Phe Gly Gln Phe Glu Leu Thr Thr Phe Asp Phe Ser Gln Tyr Met Lys Leu Asp Ile Ala Ala Val Arg Ala Leu Asn Leu Phe Gln Gly Ser Val Glu Asp Thr Thr Gly Ser Gln Ser Leu Ala Ala Leu Leu Asn Lys Cys Lys Thr Pro Gln Gly Gln Arg Leu Val Asn Gln Trp Ile Lys Gln Pro Leu Met Asp 345

Lys Asn Arg Ile Glu Glu Arg Leu Asn Leu Val Glu Ala Phe Val Glu Asp Ala Glu Leu Arg Gln Thr Leu Gln Glu Asp Leu Leu Arg Arg Phe Pro Asp Leu Asn Arg Leu Ala Lys Lys Phe Gln Arg Gln Ala Ala Asn Leu Gln Asp Cys Tyr Arg Leu Tyr Gln Gly Ile Asn Gln Leu Pro Asn Val Ile Gln Ala Leu Glu Lys His Glu Gly Lys His Gln Lys Leu Leu Leu Ala Val Phe Val Thr Pro Leu Thr Asp Leu Arg Ser Asp Phe Ser Lys Phe Gln Glu Met Ile Glu Thr Thr Leu Asp Met Asp Gln Val Glu Asn His Glu Phe Leu Val Lys Pro Ser Phe Asp Pro Asn Leu Ser Glu Leu Arg Glu Ile Met Asn Asp Leu Glu Lys Lys Met Gln Ser Thr Leu 485 Ile Ser Ala Ala Arg Asp Leu Gly Leu Asp Pro Gly Lys Gln Ile Lys 505 Leu Asp Ser Ser Ala Gln Phe Gly Tyr Tyr Phe Arg Val Thr Cys Lys Glu Glu Lys Val Leu Arg Asn Asn Lys Asn Phe Ser Thr Val Asp Ile Gln Lys Asn Gly Val Lys Phe Thr Asn Ser Lys Leu Thr Ser Leu Asn 550 555 Glu Glu Tyr Thr Lys Asn Lys Thr Glu Tyr Glu Glu Ala Gln Asp Ala Ile Val Lys Glu Ile Val Asn Ile Ser Ser Gly Tyr Val Glu Pro Met Gln Thr Leu Asn Asp Val Leu Ala Gln Leu Asp Ala Val Val Ser Phe Ala His Val Ser Asn Gly Ala Pro Val Pro Tyr Val Arg Pro Ala Ile Leu Glu Lys Gly Gin Gly Arg Ile Ile Leu Lys Ala Ser Arg His Ala 630 Cys Val Glu Val Gln Asp Glu Ile Ala Phe Ile Pro Asn Asp Val Tyr 645 Phe Glu Lys Asp Lys Gln Met Phe His Ile Ile Thr Gly Pro Asn Met 665 Gly Gly Lys Ser Thr Tyr Ile Arg Gln Thr Gly Val Ile Val Leu Met

Ala Gln Ile Gly Cys Phe Val Pro Cys Glu Ser Ala Glu Val Ser Ile 695 Val Asp Cys Ile Leu Ala Arg Val Gly Ala Gly Asp Ser Gln Leu Lys Gly Val Ser Thr Phe Met Ala Glu Met Leu Glu Thr Ala Ser Ile Leu Arg Ser Ala Thr Lys Asp Ser Leu Ile Ile Asp Glu Leu Gly Arg Gly Thr Ser Thr Tyr Asp Gly Phe Gly Leu Ala Trp Ala Ile Ser Glu Tyr Ile Ala Thr Lys Ile Gly Ala Phe Cys Met Phe Ala Thr His Phe His Glu Leu Thr Ala Leu Ala Asn Gln Ile Pro Thr Val Asn Asn Leu His Val Thr Ala Leu Thr Thr Glu Glu Thr Leu Thr Met Leu Tyr Gln Val Lys Lys Gly Val Cys Asp Gln Ser Phe Gly Ile His Val Ala Glu 825 Leu Ala Asn Phe Pro Lys His Val Ile Glu Cys Ala Lys Gln Lys Ala Leu Glu Leu Glu Glu Phe Gln Tyr Ile Gly Glu Ser Gln Gly Tyr Asp 855 Ile Met Glu Pro Ala Ala Lys Lys Cys Tyr Leu Glu Arg Glu Gln Gly Glu Lys Ile Ile Gln Glu Phe Leu Ser Lys Val Lys Gln Met Pro Phe 885 890 Thr Glu Met Ser Glu Glu Asn Ile Thr Ile Lys Leu Lys Gln Leu Lys 905 Ala Glu Val Ile Ala Lys Asn Asn Ser Phe Val Asn Glu Ile Ile Ser Arg Ile Lys Val Thr Thr 930

(2) INFORMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 23 base pairs (B) TYPE: nucleic acid

 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:	
CTGGATCCAC NGGNCCNAAY ATG	23
(2) INFORMATION FOR SEQ ID NO:4:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 23 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: cDNA	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(*i) SEQUENCE DESCRIPTION: SEQ ID NO:4:	
CTGGATCCRT ARTGNGTNRC RAA	23
(2) INFORMATION FOR SEQ ID NO:5:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: cDNA	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:	
CCACAATGGA CACTTCTGC	19
(2) INFORMATION FOR SEQ ID NO:6:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: cDNA	
(iii) HYPOTHETICAL: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:	
CACCTGTTCC ATATGTACG	19

(2) INFORMATION FOR SEQ ID NO:7:

(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: CDNA	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:7:	
AAAATGGG!	TT GCAAACATGC	20
(2) INFO	RMATION FOR SEQ ID NO:8:	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: cDNA	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:8:	
GTGATAGT	AC TCATGGCCC	19
(2) INFO	RMATION FOR SEQ ID NO:9:	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: cDNA	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: YES	
	SEQUENCE DESCRIPTION: SEQ ID NO:9:	
AGATCTTC:	IT CIGGITCGIC	20
(2) INFO	RMATION FOR SEQ ID NO:10:	
(i)	SEQUENCE CHARACTERISTICS:	

	(A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: cDNA	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:10:	
GCCAACAA	TA ATTTCTGGTG	20
(2) INFO	RMATION FOR SEQ ID NO:11:	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: cDNA	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:11:	
TGGATAAG	AA CAGAATAGAG G	23
(2) INFO	RMATION FOR SEQ ID NO:12:	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii)	MOLECULE TYPE: cDNA	
(iii)	HYPOTHETICAL: NO	
(iv)	ANTI-SENSE: NO	
(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:12:	
CCACAATG	GA CACTTCTGC	19
(2) INFO	RMATION FOR SEQ ID NO:13:	
(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:	
CACCTGTTCC ATATGTACG	19
(2) INFORMATION FOR SEQ ID NO:14:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: cDNA	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:	
AAAATGGGTT GCAAACATGC	.20
(2) INFORMATION FOR SEQ ID NO:15:	
 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
(ii) MOLECULE TYPE: CDNA	
(iii) HYPOTHETICAL: NO	
(iv) ANTI-SENSE: NO	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:	
GTGATAGTAC TCATGGCCC	19
(2) INFORMATION FOR SEQ ID NO:16:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 23 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: cDNA	
(iii) HYPOTHETICAL: NO	

WO 95/15381 PCT/US94/13805

- 46 -

(iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

GACAATAGCT TATCAATATT ACC

23

CLAIMS

- 1. An isolated and purified DNA molecule having a sequence of at least about 20 nucleotides of hMSH2, as shown in SEQ ID NO:1.
 - 2. The DNA molecule of claim 1 which is cDNA.
 - 3. The DNA molecule of claim 1 which is labeled.
- 4. The DNA molecule of claim 1 which is operably linked to a promotor sequence.
- 5. The DNA molecule of claim 4 which upon transcription produces an RNA molecule having the sequence of native hMSH2 mRNA.
- 6. An isolated and purified DNA molecule having a sequence of at least about 20 nucleotides of an hMSH2 allele found in a tumor wherein said DNA molecule contains a mutation relative to hMSH2 shown in SEQ ID NO:1.
- 7. A method of treating a person predisposed to hereditary non-polyposis colorectal cancer to prevent accumulation of somatic mutations, comprising:

administering a DNA molecule which has a sequence of at least about 20 nucleotides of hMSH2, as shown in SEQ ID NO:1, to a person having a mutation in an hMSH2 allele which predisposes the person to hereditary non-polyposis colorectal cancer, wherein said DNA molecule is sufficient to remedy the mutation in an hMSH2 allele of the person.

- 8. The method of claim 7 wherein said DNA molecule encodes an hMSH2 protein as shown in SEQ ID NO:1.
- 9. The method of claim 7 wherein said DNA molecule spans the mutation site in said allele and remedies the mutation by recombining with said mutant allele.

- 10. The method of claim 7 wherein said DNA molecule encodes a portion of hMSH2 protein which is sufficient to provide DNA mismatch repair function
- 11. A method of determining a predisposition to cancer comprising: testing a body sample of a human to ascertan the presence of a mutation in hMSH2 which affects hMSH2 expression or hMSH2 protein function, the presence of such a mutation indicating a predisposition to cancer.
 - 12. The method of claim 11 wherein the sample is DNA.
 - 13. The method of claim 11 wherein the sample is RNA.
 - 14. The method of claim 11 wherein the sample is protein.
- 15. The method of claim 11 wherein the sample is isolated from prenatal or embryonic cells.
- 16. A method of screening to identify therapeutic agents which can prevent or ameliorate tumors, comprising:

contacting a test compound with a cell;

determining the ability of the cell to perform DNA mismatch repair, a test compound which increases the ability of said cell to perform DNA mismatch repair being a potential therapeutic agent.

- 17. The method of claim 16 wherein the cell contains an hMSH2 mutation found in a tumor.
- 18. A method of screening to identify therapeutic agents which can prevent or ameliorate tumors, comprising:

contacting a test compound with an isolated and purfied hMSH2 protein;

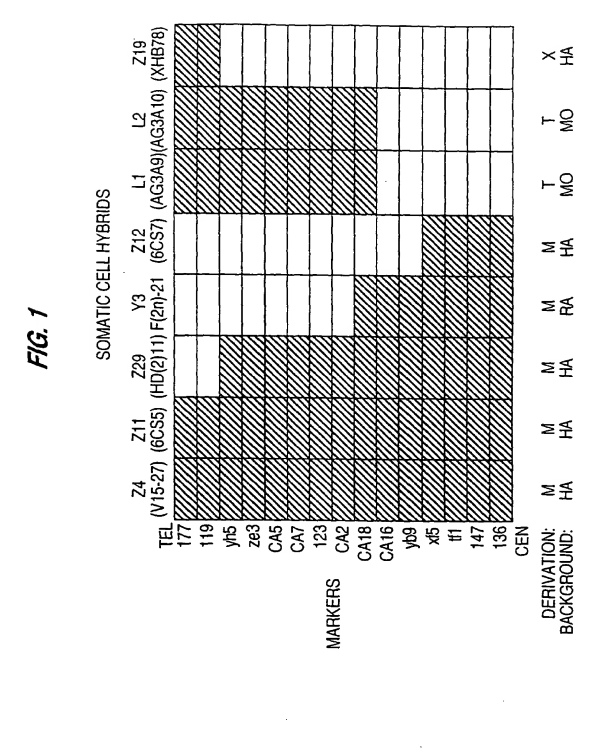
determining the ability of the hMSH2 protein to perform DNA mismatch repair, a test compound which increases the ability of said protein to perform DNA mismatch repair being a potential therapeutic agent.

19. The method of claim 18 wherein the hMSH2 protein contains a mutation found in a tumor.

- 20. An isolated and purified protein which has the sequence shown in SEQ ID NO:2.
 - 21. A transgenic animal wherein the transgene is an hMSH2 allele.
- 22. The transgenic animal of claim 21 wherein the allele is a mutant allele of hMSH2 found in a patient having hereditary nonpolyposis colorectal cancer or an RER⁺ tumor.
 - 23. A non-human animal which contains no wild-type MSH2 allele.
- 24. The transgenic animal of claim 21 wherein said animal contains no wild-type MSH2 allele.
- 25. A method of treating a person having an RER⁺ tumor to prevent accumulation of somatic mutations leading to resistance to an anti-cancer therapeutic agent, comprising:

administering to a person having a tumor with an RER⁺ phenotype: (a) a DNA molecule which has a sequence of at least about 20 nucleotides of hMSH2, as shown in SEQ ID NO:1, wherein said DNA molecule is sufficient to remedy a mutation in an hMSH2 allele of the person; and (b) an anti-neoplastic therapeutic agent.

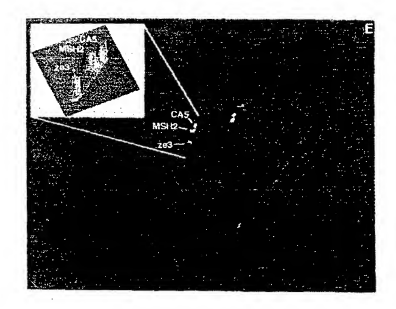
26. The method of claim 23 wherein the anti-neoplastic therapeutic agent is selected from the group consisting of: a hormone, radiation, a cytotoxic drug, a cytotoxin, and an antibody.

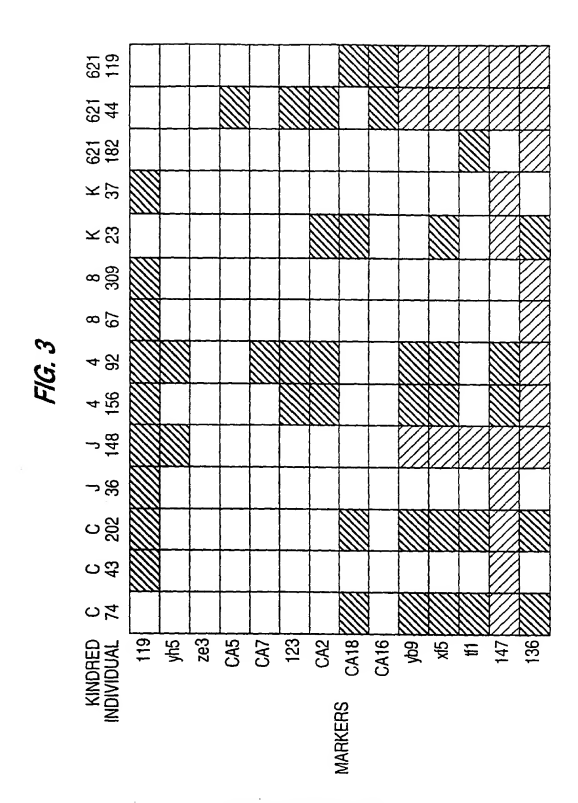


SUBSTITUTE SHEET (RULE 26)

FIG. 2A FIG. 2B FIG. 2C FIG. 2D

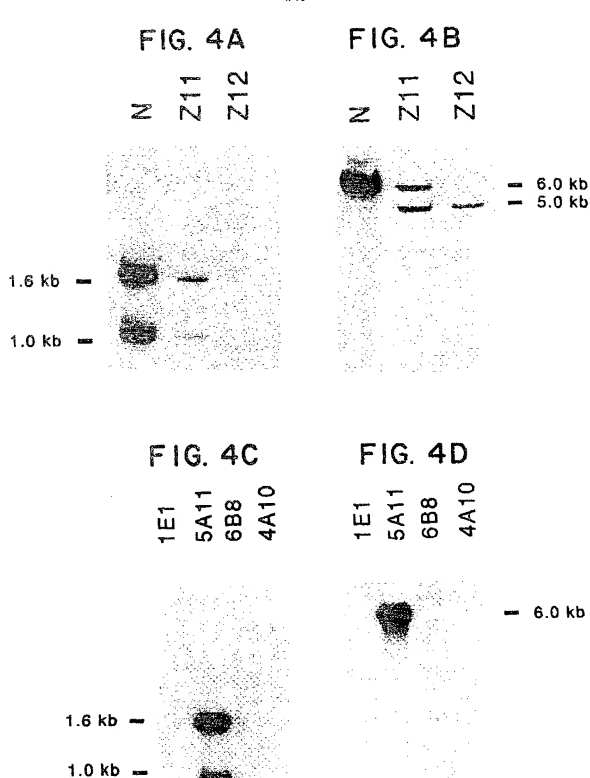
FIG. 2E





SUBSTITUTE SHEET (RULE 26)

4/10



SUBSTITUTE SHEET (RULE 26)

F16. 5,

5/10 CIA Jhr 747 ACC CAI ¥¥ Val 221 ر د د Ä ACC ιys Asn Cys 161 Ser ICI AAT Ser CAT Arg Ala Leu Asn ¥¢ Pro 040 CAG Phe CAA CAA CAA CAA Ser 2 olo CAA ASU CAI AIG Lys Pro Lys CCA AAG AAG Phe 111 AIG Val Lys X V ၁၁၁ Thr ACA Ser ASP Het 7 Cl. SAC CAG טור פענ Asn Ser CAI Ser 221 cto CAG 3 YCY **Y**1 Pro Arg CC.Y 211 122 Net Ato Ata **₹**25 Asp Asn Asn AAI ZY C 215 ב 0 CAG <u>2</u> CIC Alo Ala Val ž Ala AAC A1G Lys AAG LYS ۲۵۱ 1 c 212 פנת CAA **V**25 A I B ۲ ک GLY <u>ک</u> GIN 252 222 Ser Asn ** 155 lyr 61.7 66.4 Let Y) IAI Lys 114 clo ر دود AGG AGG TGA CAG 010 Leu ςį CCA Phe Ξ כן ל ეეე 11c 61 y 66 A Lys AAA Phe 111 = 114 Phe 111 <u>.</u> CIG CIC Val ۲I CCT ١٩٥ Asp \ \ \ 113 ٨rg **V**CV Lys AAG CAI GGC GGC Phe A L a ນນນ = 110 Arg AGA = H CTC ct CA Va V cl Y **Y**00 AIC Met Lys Leu 116 **₹**₹ Arg J Asp GAC Asn AAT Asp A1.8 GCA 111 1C1 1CA ACC Ser AGT CAC 10/ 113 11e 1.02 1.02 Vel ¥ **₹1**5 ٧aا ctu GAG o e c t c 213 c Ye Lys AAG glu CAG A 1 8 | eu AIG ეეე Ser Asn Leu (01.7 000 **Let** C16 cto CAG Phe 17r 1A1 == **V**CV Iyr Phe GIN Arg Arg CCC Ser I 101 cl Y ပ္ပ CAC Asn AAT ۲ه ا 115 c Cla CAG CAG Clu Thr Ala His Arg t eu ASO CAG CIG ₹ø^ 210 929 Lys 3 Y **Le**u C16 Ser S S ၁၁၁ 1 C C C 101 Ser Ser AGC 7 clu 292 Gin Phe CAG 11C SAG ACG Al. GCA V ... רפנ CIC Asp CAI lys AAA Asp GAC Ser Phe 721 17r 1AT S A I a ပ္ပ 01. 00.4 00.4 Arg AGA Asn YY כנץ כככ cye Cye Asp <u>ره</u> ۷ 113 CAC 787 115 Alb Phe Asp CA1 221 212 929 110 Ata GCA 17r 141 cl y ၁၅၅ Ala **₹**35 Het A1G 17. 17. A18 GCA Phe Ξ Asp GAC cto Cto Asn Sef Pro CCG Pro ວ AGC Ser)) Asp CAC 1 = E Thr \ \ \ 110 ACI CCI 1AG 1GG cl:o CAG 01 00 00 00 AIG Asp GAI 01.7 000 A59 CG1 Ser 101 ž CLY CCA glu ASP GAC CAG 핕 ACI Arg 222 Gln Leu Het A16 ۲ ۱ CCT Pro CCI ιys YYY ۲ کا در 213 V8 C Asn <u>5</u> Lys AA AAT CAG Asp 7yr 7AC CYC 300 (ys AAG Va t G11 Phe 110 ng OY **YYS** 1hr AC1 1h7 ACA ACA (Lea Phe 110 Lys 2 C11 פור Z ςlλ cyc CAG Iyr 199 CAA Ser H AIG CAA ACG C11 בפ ۲٥ SCA Cys CI Y CCA Thr C11 70/ 212 101 Phe III c U **GAA** CCC ۷ها 100 ZY. Let 21 100 G17 GGA 115 ASP GAC ر ا ه/ Y) 787 LYS AA 7 <u>≻</u> 60 00 ¥ IYI 100 Ala CC1 S 3 9 2 34 88 118 146 174 520 <u>8</u>62 258 772 5 8 8 8 286 856 202 604

F16. 5E

6/10 1¥ 1 Lec כנים **V**90 ₹ ₹ ιys ₹ CAG 010 ₹ X of ניפני Asn ¥ Thr ACC ۱ø۸ = Gla CA. ١٥٨ **9**¥3 Ser 3 ίγs ₹ Pro נור Asn Ţ ₹3 AGY CCT IYI lyr 141 **Y**55 Phe Phe II 3 <u>دا</u> < 3 ₹ Phe Asp GAC _ 5 5 Asp Arg CG1 CAG Y)) 241 Glu CAA A A 23 Lys AG A3n פוט ιys ¥G ¥ Leu 11G Leu CAA GAA 3 <u>ار</u> ه ۷۵ ا 1 - 10 Pro o U ¥ CCT Lys ¥€ **919** 001 Pro 221 Ser CCT oly ccc ASU 1<u>7</u>1 Pro CCI Val GIC \ \ \ 113 ¥ 212 Ala cye Cye 4 5 6 4 Ihr ACC ٦ø/) [Phe clo Leu CII מני ל מני A La GCA Lys ₹ Lec IIA ¥ Lys VVV 160 207 ATI TA <u>:</u> ¥I¥ Asp CAC P.e CII ე; (} 111 Asp GAT Ser ۷ء ا Ser 101 TCA CC \ 15 Asn Arg CGA Cys 101 ۷ها ۲۱۵ Ser 101 Ser Arg 1CA ?} ?} Ser **YCY** Asn Cys 161 īĥ ACT AAI AAG Leu 116 Asn Pro Lys Asn Arg CG1 CCT AAC ¥ Ala Lea ۸۱۵ 100 l ys ¥c 116 Ser 1CA AAT Asn Arg 2 **Y**CY <u>2</u> Pro CCT C11 Lys 113 ₹ Ala Asn **Y**25 Cys 161 ίγs ¥ Ξ 2 \ Va. 010 7 of c rec C 212 Asp CAG Asp **7** Let CIA 3 CIA 7 Ser Arg Thr ACC Ser YCC \ V 213 HIS AGG AGT CAG 20 Pro CCA ક 3 2 Ala Leu Th' Glu lle GAA A11 211 222 AC I <u>-</u> AIA CIA Asn \ \ \ AAC Ata GC I Ser 221 = YIY Phe 110 Asn AT 2 <u>-</u>13 Phe 110 3 IIA Arg CG1 V I O Y) ACC Ę Phe 111 Arg Ser Leu Ala 55 **Y**CY ٨rg YUU <u>=</u> YIY Pro 200 clu 3 ۲<u>۹</u> Phe II Phe 111 Lys AA Ĺγs ₹ Ser Arg Asn TCT CTG AAC 190 01.7 00.1 1 AC1 Ï CAI Ser ICA <u>ر</u>ه / Leo TA 17r 1AC Lys * G1.1 Val GIC Glu Asn Lys AAG 2 of n CAG 010 ۱ ا ا C11 cye CYe 110 = <u>ں</u> __ YIY الا 1 <u>ره /</u> AIT 18 12 13 5 CAG Asp CAT] Y Phe 11 17F 17T Fet ATG 01 00A Sl ALA Ala GCT <u>-</u> 100 AIT Hel Asp GAT **A**16 Ser 101 <u>5</u> 5 78/ 115 <u>-</u> 210 Asu Arg Lys Phe II ₹ Asp GAT Asp GAT **V**CA cl y ၁၁၁ 010 3 CIC ₹ A79 CCA A! B **₹**25 c ge of n CAG c ye Lys. AAG ¥g Leu CIA cly **Y**95 Gin Lys g K Pro CCI **A B D** of C ACT 2 116 I P ₹ 7 7 A l a **Y**33 CYC ALA GCC chg CAG Clu 3 5 **9**Y3 2 님 YCC <u>5</u> 116 Xet **Y**11 AIG 1.6 1.16 کلا در در ζ 161 Ser AGT AIC Ala cl Y **V**00 100 γsp CAT ¥¥G **11** V I D Asp GAT 5 ίγs Asp 3 ₹ Asp GAC Gln 1hr ACT 3 Ser TCC CYI 7 1 € 1 × 1 t ys ₹ 210 **CAA** = AI I <u>و</u> ک CAG Lys ₹ 3 <u>₹</u> פות Asn AAT SAC Asp GAT <u>اء</u> ۲ 17 17 **61 ده ا** 115 Ţrp 201 Arg AGG S S CAG 25 Va (Xet A1G 2 Ę 010 ב Asp GAT רפנ 116 ACT ACT ¥ c U rec o Ser 116 Asn H.S CAC 101 ΥC ౼ = **ACA** Ser Thr Asn AAT AGT ACA <u>ار</u> Asn 5 5 ¥ \$ \ 0 0 0 0 0 0 Łys ₹ פור _ 5 5 ₹ Phe AI I 111 L\s \ \ Ala Asn ₹ 1 P. Y 370 1108 398 1192 1276 1024 426 510 1528 35 485 7771 1612 588 889 594 1780

F16.50

ACA Ihr Cys 160 Ξ AC 1 Ihr. 200 ₹ Ser))] CC. √30 Leu 3 199 Cl \ Clu Val 010 ξ ון היי CAG 010 **₹**3 T o K AIG Ţyr AIG = 1 A C CAI Het Ala כאא מא AGI Ihr ACT AIG ĭ Ϋ́ C C **YY**5 Thr Acc Phe Phe Ξ 1hr AC 1 ٧a٧ CIA Pro CCA Asn 1 Ξ AAI Pro Asn ₹ Ser ICA A (B 100 Ser X : Leo 11 His CAA Pro 7 ວວວ CAI Va! CIA 101 222 Z o K AIG 1hr AC1 Thr ACC I, ACC Het Het Phe Pro Lys AAG AIG AIG CIC 111 01.y Cys 101 Phe 61 y A L B CCA 010 Clu CAG 100 11e ATC CAA Ser CCI AII AGC ۲ کن Pro Ar9 AGA Phe His Ile 11e Thr TTC CAC ATC ATT ACT Ihr T ACG CAA CAA Phe 111 Phe Asp GAI Asn 110 Ĺγš 444 AAI Val 010 C1.7 Ser Het ATG Asn 221 Thr ACT AAT 1√ 171 Asn \ V ១១ ¥. 144 215 11G Phe Ξ Ala Va (Cys 160 15r 100 Cly CGA Lys AAG Lys AAG ICA Gl 🗸 **Y**55 Cl. Cys 101 Ϋ́ Phe 111 ار د CIC رد و در Ala כו Ala Glu Leu Ser 221 CCA <u>|</u> 999 Lys ¥¥ Asp GAT At a **₽** CCA CAG Ser 160 CIG 22 AGG TAA Asp Lys Gln Het = H 91 AM CAG ATG Gln Leu teu lle 11e 11e YIY 61 y YCY ور د۲ Tr. CCA ر ده / Phe 110 ₹15 ct, ž AIC = 61.y 66.A 11 ٧٥ ا 212 - | c f c O CA 115 GIA AIG GAA IGA GAG Ala ၁၁၅ Ser ¥GT ATA Lys AAG H is X i S CYI 7 ر۷و د۲و Ala CAI AIG Asp Ĭ S 114 Ale Thr GCA ACA ر د] c CIA A11 1yr ۳ H Lys AAA IAI Glu Lys GAA AAA Ser ICA Lec CIC ci 🗸 200 Asn Clu AAI ci y 999 == CAG AIT <u>2</u> ۲<u>۱</u>۷ CIA Ala 133 Asp GAT Yo. = Asn YY1 AII Phe Ξ Phe 111 ίγs ¥¥ Sla CAG Y)) Phe III == AIA 999 Lys cl y 1,7 1,7 0 6 6 6 7°7 Ser AGT S S Lys YYY AAA ATC 7. ∑. GTG Clu 700 Val 715 ٦ ე ე ე \ \ \ ACC 1hr Ac1 **₹** CAG <u>5</u> rec C 1 Asn Asp Val Alb 61 y 666 Arg CGA Pro CCA **V**25 Ser ICA Asp C11 CAI 5 ₹ Lys 75 Leu Glu Leu ۲ ၁၁၅ Ser AIA Thr AIA 101 = 0 PC C ACG 101 = ٧Y At a Glu 3 t eu Arg AGG c C C \ \ \ \ 219 213 AGA īĥ ACA ACT Arg CGA Ala CCT 17p Asn 155 250 225 ₹ == AIC 611 11e A I & 201 Y) Ata GCC L ys Lys ₹ Gln Ser Lec 74 בים בים Lys 101 AAG IVI Z 1948 846 2536 734 762 2284 790 818 2452 2872 2368 874 2620 2704 SUBSTITUTE SHEET (RULE 26)

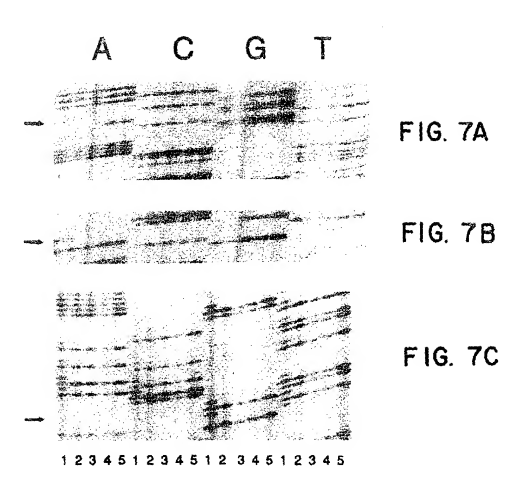
7/10

F16. 64

	80 80	160	156	236	536 236	316	311	379	391	453	470	532
SH2 SH2 SH2 SH2 SH2 SH2 SH2 SH2 SH2							Weudinkingddlals	Vedittesqs[f]A		L [erf ee ln <mark>reaktforo</mark> aa <mark>wkoe</mark> cy e lkoginqlewkoalekhegkhak		HENT LIND ON BALLE- FINAR PSER PARELLE FINAR FOR THE SARROTS LISAR BOOK OF BYYFRAMER OF THE PROPERTY.
출축 결국 결국 결국 결국 기업	hMSH2 yMSH2	hм SH2	умѕн2	hMSH2	yMSH2	hMSH2	yMSH2	hmsh2	умѕнг	hMSH2	умѕн2	hMSH2

F1G. 6B

умѕн2	<u>Vett</u> ivologa yeenneen koen eekkinskoot oo kaan oo kaan oo kaa oo kaan kaan hamaan oo kaan oo	550
hмsн2	INNINASTVDIORNEVERNSKOTSLNESYTKNKTEVERAODAIVKENNIISSGIVEPMOTONDIINDIIRAOIODAIVSENINS	612
yMSH2	<u>inakitkav</u> ielstvora <u>en</u> festkoloksianetniloke <u>nvo</u> koosolu <u>krettinist</u> ltytpvfekosl <u>utahind</u> voastanes	630
SOS PWSH2	NGARVPYVRPailekggg-EiiEkasreacvevodelatarpanyrfekokomghitvernwegkstvirotavivlamon	691
STITUTE STUTIES	SY <u>APITPYIRP</u> KIhpmdserRthLis <u>srh</u> PvlDMODDIspisms <u>NDV</u> TLESGKGDFLITWGDNWGGRSTVIROVGVGVISLAMODI	710
SHEE SHEE	GGEVPCESAEVSIVDCILVARVGAGDSOLKGVSTFWADMLETASILRSATKDSLITIDELGRGTSTYDGFGLAWAISEYITA	771
J (RULE	GCFVPCEBABIAKVDALLCRVGAGDSOLKGVSTFMVEI [DBTAS)LLKMASKNSLITIVDELGRGTSTYDGFGLAWAI AEHITA	9/10 ₀ 6/
26)	<u>Innigafompanhehen</u> alangiPnim <u>nihi</u> talee <u>ep</u> tl <u>imin</u> oukk <u>enoberhinae</u> lam <u>er</u> hi <u>nf</u> e	842
умѕн2	SKIGCFAIDDAYHIBHBIY E L SEKL PMUKNWHVA hieknlkeqkhd <u>DB</u> DI WBAY KVEP GI S DOSEGIHVA BVVQBBK IV K	870
hMSH2	CANORALIBBISEQYIGESqgydi n epaakkcyle d eqgekiiqefls N okqmoftemsegni d iklkglkaev la knnsf	922
умѕн2	wakkanaanouktundoookakuslqevnegnigikalikewirkvBeegOhdpskitaeagqhkigeliraEenek	950
hMSH2	v <mark>ne</mark> iisr ii kv i t	934
yMSH2	e <u>ne</u> ny i eliy k <u>s</u> pocyn	966



			•
		./	
, - , ·	·		